Chemical characteristics of size resolved atmospheric aerosols in Iasi, north-eastern Romania: Nitrogen-containing inorganic compounds control aerosols chemistry in the area

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Abstract. This study assesses the effects of particle size and season on the content of the major inorganic and organic aerosol ionic components in the Iasi urban area, north-eastern Romania. Continuous measurements were carried out over 2016 by means of a cascade Dekati Low-Pressure Impactor (DLPI) performing aerosol size classification in 13 specific fractions over the 0.0276-9.94 µm size range. Fine particulate Cl−, NO3−, NH4+ and K+ exhibited clear minima during the warm seasons and clear maxima over the cold seasons, mainly due to trends in emission sources, changes in the mixing layer depth and specific meteorological conditions. Fine particulate SO42− did not show much variation with respect to seasons.

Particulate NH4+ and NO3− ions were identified as critical parameters controlling aerosols chemistry in the area, and their measured concentrations in fine mode (PM2.5) aerosols were found to be in reasonable good agreement with modelled values for winter but not for summer. The likely reason is that NH4NO3 aerosols are lost due to volatility over the warm seasons. We found that NH4+ in PM2.5 is primarily associated with SO42− and NO3− but not with Cl−. Actually, indirect ISORROPIA-II estimations showed that the atmosphere in the Iasi area might be ammonia-rich during both the cold and warm seasons, enabling enough NH3 to be present to neutralize H2SO4, HNO3 and HCl acidic components and to generate fine particulate ammonium salts, in the form of (NH4)2SO4, NH4NO3 and NH4Cl. ISORROPIA-II runs allowed us to estimate that over the warm seasons ~ 35 % of the total analysed samples had very strong acidic pH (0–3), a fraction that rose to ~ 43 % over the cold seasons. Moreover, while in the cold seasons the acidity is mainly accounted for by inorganic acids, in the warm ones there is an important contribution by other compounds, possibly organic. Indeed, changes in aerosols acidity would most likely impact the gas–particle partitioning of semi-volatile organic acids. Overall, we estimate that within the aerosol mass concentration the ionic mass brings contribution as high as 40.6 %, with the rest still being unaccounted for.

Keywords: urban aerosols, ionic chemical composition, ammonium, nitrate, pH, long range transport, Iasi, Romania.
1 Introduction

Despite dramatic progress made to improve air quality, global air pollution continues harming people’s health and the environment. The problem of aerosols or particulate matter (PM) is still a great concern (Olariu et al., 2015) in Europe and many other areas in the world (e.g., China, India, USA). Atmospheric aerosols, described as complex mixtures of liquid and/or solid particles suspended in a gas (Olariu et al., 2015), are mainly originating from anthropogenic and natural sources (Querol et al., 2004). Fine PM$_{2.5}$ particles (airborne particles with an equivalent aerodynamic diameter < 2.5 µm) are air pollutants with significant effects on human health (Pope et al., 2004; Dominici et al., 2006; WHO, 2006a; Directive 2008/50/EC, 2008; Sicard et al., 2011; Ostro et al., 2014) as well as air quality (Directive 2008/50/EC, 2008; Freney et al., 2014), visibility (Tsai and Cheng, 1999; Directive 2008/50/EC, 2008), ecosystems, weather and climate (Ramanathan et al., 2001; Directive 2008/50/EC, 2008; IPCC, 2013). Aerosols are also known to play a significant role within the chemistry of the atmosphere (Prinn, 2003), acting as surfaces for heterogeneous chemical reactions (Ravishankara, 1997).

Safety threshold values for both PM$_{2.5}$ (10 or 17 µg m$^{-3}$ air, as annual mean) and PM$_{10}$ (airborne particles with an equivalent aerodynamic diameter < 10 µm; 20 or 28 µg m$^{-3}$ air, as annual mean) are addressed by the WHO (2006b) or by the 2008/50/EC Directive (2008) on ambient air quality and cleaner Europe. With regard to the PM$_{2.5}$ fraction, the EEA Report (2015) indicates that in 2013 the EU daily limit values for PM$_{10}$ and PM$_{2.5}$ were exceeded in, respectively, 22 and 7 out of the 28 EU member states. However, a decreasing trend was observed when compared with the WHO Report data (2006b). Moreover, on a global scale the PM$_{2.5}$ exposure leads to about 3.3 million premature deaths per year (predominantly in Asia), a figure that could double by 2050 (Lelieveld et al., 2015). Indeed, PM air pollution imparts a tremendous burden to the global public health, because it ranks as the 13th leading cause of mortality (Brook, 2008).

Up to now the chemical composition of atmospheric aerosols is reported for various urban European sites (Bardouki et al., 2003; Hitzenberger et al., 2006; Tursic et al., 2006; Gerasopoulos et al., 2007; Schwarz et al., 2012; Laongsri and Harrison, 2013; Wonaschutz et al., 2015; Sandrini et al., 2016), but such information is very scarce in Eastern Europe (Arsene et al., 2011). Sulfate (SO$_4^{2-}$), nitrate (NO$_3^-$) and ammonium (NH$_4^+$) ions, which are major inorganic particle constituents (Wang et al., 2005; Bressi et al., 2013; Hasheminassab et al., 2014; Voutsa et al., 2014) are mainly secondary species formed in the atmosphere by chemical reactions of their precursor gases and by physical processes (nucleation, condensation, etc.) (Aksoyoglu et al., 2017). Ammonium aerosols, with an atmospheric lifetime of 1–15 days, have a clear tendency to deposit at large distances from their emission sources (Aneja et al., 2000) and seem to play a very important role in atmospheric chemistry. In urban air, the abundance of NO$_3^-$ on fine particles seems to mainly depend on the reaction between HNO$_3$ and NH$_3$ (Stockwell et al., 2000). On a global scale, HNO$_3$ heterogeneous reactions on mineral dust and sea salt particles might be the predominant source of particulate NO$_3^-$ (Athanasopoulou et al., 2008; Karydis et al., 2011). The human health effects of atmospheric ammonia, primarily exerted through particulate NH$_4$NO$_3$, are gradually acquiring importance compared to NO$_x$ emissions (Sutton et al., 2011).
Relative humidity (RH; Jang et al., 2002) and gas–particle partitioning processes of semi-volatile species (NH$_3$, HNO$_3$, HCl and some organic acids) affect the hydronium ion (H$_3$O$^+$) concentrations (Keene et al., 2004; Guo et al., 2016) as well as aerosol formation and chemical composition. Aerosol acidity is often estimated with ion balance methods (Trebs et al., 2005; Metzger et al., 2006; Arsene et al., 2011), phase partitioning (Meskhidze et al., 2003; Keene et al., 2004) and thermodynamic equilibrium models (e.g., E-AIM, Clegg et al., 1998; Wexler and Clegg, 2002) or ISORROPIA-II (Nenes et al., 1999; Fountoukis and Nenes, 2007; Hennigan et al., 2015; Guo et al., 2015, 2016; Fang et al., 2017). Acidity might influence transition metals solubility and enhance aerosols toxicity and the delivery of nutrients by atmospheric deposition in marine areas (Meskhidze et al., 2003; Fang et al., 2017).

Unfortunately, the aerosols role in the global atmospheric system is not yet sufficiently understood. The main related challenges are the occurrence of multiple sources (e.g., soil erosion, sea spray, biogenic emissions, volcano eruptions, soot from combustion, condensation of precursor gases) and the complexity of interactions with other atmospheric constituents (Zhang et al., 2015). Nowadays, sources, distribution and behaviour of natural and anthropogenic aerosols are still a matter of debate, exacerbate by the scarcity of aerosols-related work for eastern EU countries (EEA Report 5, 2015) but also by the existent discrepancies between models and field measurements. The main uncertainties are related to secondary inorganic aerosols that control the availability of atmospheric sulfuric acid (H$_2$SO$_4$), nitric acid (HNO$_3$) and ammonia (NH$_3$) (Ianniello et al., 2011). The existing knowledge gaps bring high uncertainty in the estimated radiative forcing, although they do not impair the conclusion that warming of the climate system is unequivocal (IPCC, 2013).

Despite a growing international recognition of the importance of air pollution and air quality problems, there is a definite need to assess air pollution patterns in Romania. The existing data for north-eastern Romania concern the chemical characteristics of ambient air pollutants such as the water-soluble ionic constituents of aerosols (Arsene et al., 2011) and rainwater (Arsene et al., 2007). Recent work performed by Arsene’s group in the field of atmospheric chemistry has clearly shown that, in the Iasi urban environment (north-eastern Romania), ~ 59 % of the total aerosol mass concentration is still unaccounted for. The present work reports for the first time detailed information on the chemical composition and seasonal variation of size-segregated, water-soluble ions in aerosol samples collected throughout 2016 in the Iasi urban area, also taking into account the potential ongoing chemistry and the contributions of critical driving forces such as meteorological factors (relative humidity, temperature), mixing layer depth and emission sources intensity. As a first attempt to assess particles acidity in the area, the present work highlights the existence of significant aerosol fractions characterized by pH values in the very strong acidity range (0-3 pH units), with potentially important implications on acid rain. Moreover, the potential importance of gaseous precursors (e.g., NH$_3$, HNO$_3$, HCl) on secondary inorganic PM is also discussed.
2 Experimental

2.1 Measurement site

Measurements were performed in Iasi, north-eastern Romania, at the Air Quality Monitoring Station (AMOS, 47°9' N latitude and 27°35' E longitude) of the Integrated Centre of Environmental Science Studies in the North Eastern Region, "Alexandru Ioan Cuza" University of Iasi, CERNESIM-UAIC, Romania. AMOS is located north-east from the city centre, on the rooftop of the highest University building (~35 m above the ground level), in a totally open area that characterises the site as a urban receptor point most probably influenced by well-mixed air masses. A comprehensive demo-geographical characterization of Iasi is described in detail by Arsene et al. (2007, 2011). However, according to a more recent estimate of the Romanian National Institute of Statistics, in 2016 the population in Iasi reached about 362,142 inhabitants (Ichim et al., 2016).

2.2 Field measurements

Size resolved atmospheric aerosols were collected on ungreased aluminium filters (25 mm diameter) using a cascade Dekati Low-Pressure Impactor (DLPI) operating at a flow rate of 29.85 L min⁻¹. Similar devices have been successfully used in other studies (Kocak et al., 2007; Wonaschutz et al., 2015). The DLPI unit performs aerosol size classification in 13 specific fractions (with size cuts at 0.0276, 0.0556, 0.0945, 0.155, 0.260, 0.381, 0.612, 0.946, 1.60, 2.39, 3.99, 6.58 and 9.94 µm at 50 % calibrated aerodynamic cut-point diameters and at 21.7 °C, inlet pressure 1013.3 mbar, outlet pressure 100 mbar and 29.85 L min⁻¹ flow rate). Sampling performance of the DLPI unit was verified through comparison with simultaneous measurements performed with a stacked filter unit (SFU) system previously used by Arsene et al. (2011). Before each reuse the sampler’s components were cleaned with ultra pure water and methanol. Disposable polyethylene gloves were always used to avoid hand contact with sampler's components, which were assembled and disassembled in a Labguard Class II Safety Biological Cabinet, NuAire. The DLPI sampler was transported to and from the field in tightened polyethylene bags. Sampling was performed in 2016 twice a week, on weekend and working days, for a total of 84 sampling events (41 during the cold seasons from October to March, and 43 during the warm seasons from April to September), generating 1092 size resolved aerosol samples. Sampling took place on a 36 h basis, with each sampling event starting at 18:00 local time. It was collected an average volume of 64.33 ± 0.85 m³ per sampling. At least two field blanks (consisting of loaded sampler taken to and from the field but never removed from its tightened bag) were generated and simultaneously analysed together with the laboratory blanks, in order to assess possible contamination during sampler loading, transport, or analysis. Meteorological parameters including atmospheric temperature (AT), relative humidity (RH), wind speed (WS), wind direction and global radiation were provided by a Hawk GSM-240 weather station, running at the AMOS site. Information about the mixing layer depth (atmospheric boundary layer) and its ability to dilute atmospheric pollutants at the investigated site was obtained from the NOAA Air Resources Laboratory (ARL) website (Stein et al., 2015; Rolph et al., 2017).
2.3 Sample analyses

Aerosols masses for both the PM$_{2.5}$ and PM$_{10}$ fractions were gravimetrically determined with a Sartorius microbalance (MSU2.7S-000-DF, ± 0.2 µg sensitivity) by weighing aluminium filters before and after sampling. Before weighing, filters stored in Petri dishes were kept for at least 3 days in a conditioned room at 40 ± 2 % RH and at 20 ± 2 °C. In between different procedure, they were appropriately stored in zipped plastic bags.

Chemical analyses for water-soluble anions (i.e., acetate, (C$_2$H$_3$O$_2$)$^-$, formate, (HCO$_2$)$^-$, fluoride, (F$^-)$, chloride, (Cl$^-)$, nitrite, (NO$_2^-$), nitrate, (NO$_3^-$), phosphate, (PO$_4^{3-}$), sulfate, (SO$_4^{2-}$) and oxalate, (C$_2$O$_4^{2-}$)) and water-soluble cations (i.e., sodium, (Na$^+$), potassium, (K$^+$), ammonium, (NH$_4^+$), magnesium, (Mg$^{2+}$) and calcium (Ca$^{2+}$)) were performed by ion chromatography (IC) on an ICS-5000$^+$ DP DIONEX system (Thermo Scientific, USA). Analyses were performed within a week from sampling, in aqueous extracts of the collected samples. For chemical analysis, steps in the analytical procedures were strictly quality-controlled to avoid contamination.

After sampling and all other required preparative steps, one half of each collected filter was ultrasonically extracted for 45 minutes in 5 mL deionized water (resistivity of 18.2 MΩ.cm) produced by a Milli-Q Advantage A10 system (Millipore). Filtered extracts (0.2 µm pore size cellulose acetate filters, ADVANTEC) were analysed on an IonPac CS12A (4×250 mm) analytical column for cations and an IonPac AS22 (4×250 mm) column for anions, running simultaneously on the IC system. The chromatographic instrumental set-up was completed by CSRS 300×4 mm and AERS 500×4 mm electrochemical suppressors and conductivity detectors. Ions analyses were performed under isocratic elution mode, using CO$_3^{2-}$/HCO$_3^-$ (4.5/1.4 mM, 1.2 mL min$^{-1}$) as mobile phase for anions and methanesulfonic acid (20 mM, 1.0 mL min$^{-1}$) for cations. Traceable standard solutions (Dionex Seven Anions II and Dionex Six Cations II) were used to generate calibration curves for each species of interest, all having correlation coefficients $R^2$ well above 0.995. The detection limits of the major species (defined as 3 times the standard deviation of blank measurements relative to the methods sensitivity) on a 36 h measurement period were: 0.0003 µg m$^{-3}$ for NH$_4^+$ (3.2 µg L$^{-1}$), 0.001 µg m$^{-3}$ for Na$^+$ (13.4 µg L$^{-1}$), 0.0015 µg m$^{-3}$ for K$^+$ (19.8 µg L$^{-1}$), 0.0002 µg m$^{-3}$ for Mg$^{2+}$ (3.2 µg L$^{-1}$), 0.0016 µg m$^{-3}$ for Ca$^{2+}$ (20.6 µg L$^{-1}$), 0.0010 µg m$^{-3}$ for SO$_4^{2-}$ (13.2 µg L$^{-1}$), 0.0012 µg m$^{-3}$ for NO$_3^-$ (15.1 µg L$^{-1}$), and 0.0019 µg m$^{-3}$ for Cl$^-$ (24.9 µg L$^{-1}$). Ions concentrations in analysed blank filters (laboratory and field) were subtracted from their corresponding concentrations in the aerosol samples.

The sum of the detected ions, or of the gravimetrically determined mass concentration, over all DLPI stages is hereafter termed “PM$_{10}$ fraction”, while the sum over impactor stages from 1 to 10 is termed “PM$_{2.5}$ fraction”. Modal diameters of the size segregated aerosols particles or of the analysed ionic components (individual or as a sum) were determined by fitting lognormal distributions.

2.4 Estimation of the aerosols acidity

The thermodynamic model proposed by Fountoukis and Nenes (2007), i.e. ISORROPIA-II (http://isorropia.eas.gatech.edu/), was used to get an estimate of the in-situ potential acidity of our PM$_{2.5}$ fractions. ISORROPIA-II thermodynamic equilibrium
model calculates the gas/liquid/solid equilibrium partitioning of $K^+$, $Ca^{2+}$, $Mg^{2+}$, $NH_4^+$, $Na^+$, $SO_4^{2-}$, $NO_3^-$, $Cl^-$ and aerosol water content, and it can predict particles pH. Up to now the model has been used in various field campaigns data analysis (Nowak et al., 2006; Fountoukis et al., 2009).

To obtain the best predictions of aerosols pH, ISORROPIA-II was run in the “forward mode” for metastable aerosol state, as preliminary runs of the experimental data in the “reverse mode” did not supply suitable information. In the “metastable” mode the aerosol is assumed to be present only in the aqueous phase, either supersaturated or not (Fountoukis and Nenes, 2007). As model input data we used just aerosol-phase ion concentrations measured by IC, along with relative humidity and temperature data from the Hawk GSM-240 weather station. Actually, in the absence of accompanying gas-phase data required to constrain the thermodynamic models, the accuracy of aerosol pH predictions can be enhanced by using the aerosol concentrations in “forward mode” calculations (Guo et al., 2015; Hennigan et al., 2015), which seem to be less sensitive to measurement errors than the “reverse mode”. Where required, NH$_3$ data predicted by ISORROPIA-II were used for the interpretation of the results.

2.5 Air mass back trajectories and air mass origin

Air mass back trajectories were calculated using the HYSPLIT 4 model of the NOAA Air Resources Laboratory (Stein et al., 2015; Rolph et al., 2017). Forty-eight hour back trajectories, arriving at the investigated site at 18:00 local time (15:00 UTC), were computed at 500, 1000 and 2000 m altitudes above the ground level. Four major sectors of air masses origin were distinguished, and their contributions are shown in FIG. 1. The most and least frequent sectors were the north-eastern (N-E, 36.6 %) and the south-south-eastern (S-SE, 11.4 %) ones, respectively. The W-SW sector is mainly prevailing during winter, while the N-E sector is most common during summer. The NW sector had a slightly enhanced frequency in winter and summer that according to James (2007), could reflect a possible European monsoon circulation. Events from the S-SE sector, prevailing mainly in spring, carried out marine chemical features highly influenced by the Black Sea. Air masses undertaking faster vertical transport most probably due to the locally/continentally driven buoyancy (Holton, 1979; Seinfeld and Pandis, 1998), travelling above large (long range transport) or short (local) continental areas were also indentified. For instance, in April 2016 five sampled events out of a total of eight were highly influenced by fast vertical air mass transport. In these events, air masses from both 500 and 1000 m altitude went down to below 500 meters (brushing the ground surface), with strong impact on the chemical composition of the collected particles (vide infra).

3 Results and discussion

3.1 Variability of PM$_{10}$ and PM$_{2.5}$ mass concentrations

Table 1 shows summary statistics (median, geometric mean, arithmetic mean, standard deviation, minimum and maximum) for PM$_{10}$ and PM$_{2.5}$ mass concentrations at the AMOS site for both working days and weekends. Statistical tests were applied to determine whether there are significant differences among working days and weekends. The Shapiro-Wilk normality test
applied to both PM$_{2.5}$ and PM$_{10}$ indicated that the entire data-base was normally distributed (detailed statistics are given in Table S 1 from Supplement Material, SM). Moreover, the difference in the mean values of the two groups was not high enough to exclude random sampling variability, thereby suggesting that the differences are not statistically significant. Therefore, local anthropogenic activities seem to bring similar contribution to the aerosol burden in the area on both working days and weekends. However, the PM$_{10}$ annual mean mass concentration (18.95 µg m$^{-3}$) did not exceed the WHO 20 µg m$^{-3}$ air annual mean value, while the PM$_{2.5}$ annual mean mass concentration (16.92 µg m$^{-3}$) exceeded the 10 µg m$^{-3}$ air annual mean value set by WHO (2006b).

Table 2 presents the annual and/or seasonal arithmetic means of PM$_{10}$ and PM$_{2.5}$ mass concentrations in Iasi, compared to other European sites (mean ± stdev). The annual averages obtained in the present work show differences in comparison with those reported by Arsene et al. (2011) for the same site. Arsene et al. (2011) have used a SFU system consisting of a 8.0 µm pore size, 47-mm diameter Isopore polycarbonate filter mounted in front of a 0.4 µm pore size, 47-mm diameter Isopore filter. However, the values determined in the present work for the fractions PM$_{0.027-1.6}$ (15.6 ± 8.7 µg m$^{-3}$) and PM$_{0.381-1.6}$ (9.1 ± 5.6 µg m$^{-3}$) are much closer to those reported by Arsene et al. (2011) for PM$_{1.5}$ (10.5 ± 11.2 µg m$^{-3}$). The potential influence on the PM levels of particle size cut-off, differences in sampling site altitude, occurrence of precipitation events, long-range transport phenomena, and air masses buoyancy is presented in detail in Section S 1 of the SM.

FIGURE 2 shows monthly arithmetic mean mass concentrations (± standard deviations) of PM$_{10}$ and PM$_{2.5}$ in Iasi. The distribution of the PM$_{2.5}$/PM$_{10}$ ratio is presented in the same figure. The relative contribution of PM$_{2.5}$ toward PM$_{10}$ showed low variability amongst the months of the year, with ratios ranging from ~ 0.75 to ~ 1.0. For urban background and/or traffic sites, WHO (2006a) suggests PM$_{2.5}$/PM$_{10}$ ratios in the 0.42–0.82 range. Seasonal patterns, with maxima during the cold and minima during the warm seasons, are observed for all of the PM$_{2.5}$, PM$_{10}$ and PM$_{2.5}$/PM$_{10}$ profiles and might be the combined result of seasonal emissions variations, local- and long-range air transport and dispersion, chemical processes, and deposition (Wang et al., 2016). The maxima of PM$_{2.5}$/PM$_{10}$ during the cold seasons are most probably caused by combustion processes as the burning of coal/petroleum for heating purposes enhances secondary aerosols generation (Li et al., 2012). The lower PM$_{2.5}$/PM$_{10}$ values during the warm seasons might be due to dust events and to more intense anthropogenic activities near the sampling site (e.g., excavation, construction and building renewal) that would both cause higher loading of coarse particles in the atmosphere. As suggested by Zhang et al. (2001), various land use categories (e.g., grass, crops, mixed farming, shrubs) corroborated with other particle-related characteristics (i.e., particle density, relevant meteorological variables) may enhance the dry deposition of sub-micron particles during the warm seasons and hence their fine/coarse ratio. Another clear seasonal pattern was observed for the mass concentration size distribution (FIG. 3). Over the cold seasons, it had a clear monomodal feature with maximum at 381 nm. In contrast, the warm seasons were characterized by the same dominant fine mode, but also by the occurrence of a super-micron mode between 1.60–2.39 µm. Again, changes in sources contributions and meteorological conditions could account for the observed differences (details in Section S 2 of the SM).
3.2 Ionic balance, seasonality of water soluble ions and stoichiometry of (NH₄)₂SO₄ and NH₄NO₃

3.2.1 Ionic balance and potential aerosols acidity

The completeness of the ionic balance was checked for the identified and quantified species (F⁻, Cl⁻, NO₂⁻, NO₃⁻, PO₄³⁻, SO₄²⁻, HCO₂⁻, C₂H₃O₂⁻, C₂O₄²⁻, Na⁺, K⁺, NH₄⁺, Mg²⁺ and Ca²⁺) in both PM₁₀ and PM₂.₅. The slopes in the raw ion chromatography data related to \( \sum \) cations vs. \( \sum \) anions were < 1 in both PM₂.₅ and PM₁₀ (detailed statistics in Table S 2), pointing to an important cation deficit that was higher in the cold compared to the warm seasons. However, for each sampling event, either cation or anion deficit was observed in various impactor stages. It should also be noted that, at the investigated site, it was even observed RH ~ 82 % during the cold seasons.

Predicting pH is suggested as the best method to analyse particle acidity (Guo et al., 2015). The ion balance method is usually based upon the principle of electroneutrality, and any deficit in measured cationic compared to anionic charge is assigned to the presence of unmeasured protons (H⁺). The reverse occurs for unmeasured hydroxyl (OH⁻) (Hennigan et al., 2015) or bicarbonate/carbonate (HCO₃⁻/CO₃²⁻) (Fountoukis and Nenes, 2007). In the present work, HCO₃⁻/CO₃²⁻ was assigned as the missing anion (details in Section S 3 for HCO₃⁻/CO₃²⁻ estimation) while NH₄⁺ as the main missing cation (details in Section S 4 for the missing NH₄⁺ assumptions). Detailed statistics of the \( \sum \) cations vs. \( \sum \) anions dependences, with HCO₃⁻/CO₃²⁻ and missing NH₄⁺ included in the ionic balance, is presented in Table S 3.

In an attempt to investigate whether or not the H⁺ species would bring an important contribution within the ionic balance, ISORROPIA-II thermodynamic equilibrium model proposed by Fountoukis and Nenes (2007) has been used in the present work (more details about ISORROPIA-II runs in Section S 5 of the SM). We investigated the relationship between ISORROPIA-II predicted aerosol pH and the ionic balance for the present data-base, and the results are presented in FIG. 4a.

The data remarkably follow a traditional titration curve, and they also clearly show that many of the analysed particles were ~ neutral (dashed lines at 0). A very important fraction of the investigated samples was in the acidic range (pH < 3 if samples were in cation deficit mode), while the remaining fraction was alkaline (pH slightly above 7 with an anion deficit mode). However, as suggested by Hennigan et al. (2015), small uncertainties in the ionic balance (mainly due to measurement uncertainties and more likely in ~ neutral conditions) may lead to shifts that span over about 10 pH units. Moreover, the sensitivity to changes in the aerosol NH₄⁺ concentration has been checked in predicted aerosol pH under forward-mode calculation. As shown in FIG. 4b, it seems that the predicted pH might decrease by 2 % when the ionic balance takes into account NH₄⁺(total) concentration (defined as the sum between that derived from raw IC data and the part estimated by using the rationale of Arsene et al. (2011)). Details on the pH sensitivity tests toward NH₄⁺ concentrations are given in Section S 6. In Iasi, north-eastern Romania, an important fraction of the total analysed samples was alkaline while the remaining part was acidic. A more detailed view of the samples pH distribution can be obtained from the data presented in FIGS. 4c,d. It seems that over the warm seasons about 55–56 % of the analysed samples were alkaline (pH > 7) and about 44–45 % were acidic (pH < 7), with the last fraction mainly distributed in the very strong acidity fraction (~ 35 % of the samples with pH in the 0–3 range, and about 2 % with aerosol pH less than 0). Over the cold seasons only 47 % of the total analysed samples were...
alkaline (pH > 7) and 53 % were acidic (pH < 7). The acidity was also mainly distributed in the very strong acidity fraction (~ 43 % of the acidic samples with pH in the 0–3 range). Sulfuric, nitric, hydrochloric and formic acids are the most likely contributors to aerosols pH in the 0–3 range. Note that strongly acidic aerosols affect air quality, health of aquatic and terrestrial ecosystems (especially through acid deposition), as well as atmospheric visibility and climate (Dockery et al., 1996; Gwynn et al., 2000; Hennigan et al., 2015). Possible impacts of strongly acidic aerosols are presented in more detail in Section S 7. Moreover, aerosols acidity can impact the gas–particle partitioning of semi-volatile organic acids. While under strongly acidic conditions (pH 1–3) the pH contribution of organic acids is expected to be negligible as these conditions prevent their dissociation, the scenario may change completely at pH values of 3–7 (vide infra). Under these circumstances, formic acid with pK_a = 3.75 (Bacarella et al., 1955) might give a significant pH contribution to ~7% of the warm-season samples and ~ 10% of the cold-season ones.

FIGURES 5a,b present the size distribution of averaged aerosol mass, NO_3^−, SO_4^{2−} and NH_4^+ concentrations while FIGS. 5c,d present ISORROPIA-II estimates for pH and H^+ mass concentration distributions, for both the cold (FIGS. 5a,c) and the warm (FIGS. 5b,d) seasons. Clear monomodal distribution seems to be specific for the cold seasons, while for the warm seasons the second mode (1.60–2.39 µm) mass concentration distribution seems to be predominated by NO_3^−. For the 155–612 nm size range, from details presented in FIGS. 5c,d, it is quite clear that pH ≤ 2. In the present work, the aerosol H^+ levels inferred indirectly from the ion balance as proposed by Hennigan et al. (2015) showed statistically significant correlation with the H^+ loadings predicted by ISORROPIA-II in the forward mode (Pearson coefficient of 0.72, p < 0.001). However, despite the good correlation, there were important discrepancies between the two estimated H^+ levels (intended as absolute values), with those from ISORROPIA-II being considerably lower than those from the ionic balance. The main issue with the model is that it may account only for partial dissociation, while the ionic balance may be affected by the uncertainty due to the propagation of measurement error. The latter may be particularly important in the presence of a slight anion deficit balance, which is interpreted as an H^+ loaded system. Despite the uncertainties in the actual pH values, it is very likely that the 155–612 nm aerosol particles are strongly acidic and that H^+ is mainly contributed by completely dissociated strong acids such as H_2SO_4 and HNO_3. Contributions from free acidity (dissociated H^+) or total acidity (free H^+ and undissociated H^+ bound to weak acids) is expected to be more important in all other remaining fraction, and especially in the 27.6–94.5 nm particles size range (vide infra). Of course, a higher confidence in the estimate of particles pH would allow better prediction of the chemical behaviour of organics that, if dissociated at relatively low acidities, would significantly contribute to the ion balance.

In the literature there is suggestion that the molar ratio approach may be a proxy for aerosol pH estimation (Hennigan et al., 2015). However, such a procedure is highly susceptible to bias the results, either due to exclusion of minor ionic species or because it does not consider the effects of aerosol water or species activities on particle acidity. In this work, even when the aerosol was inferred to be highly acidic (samples with molar ratio NH_4^+/((Cl^- + NO_3^- + 2×SO_4^{2−}) < 0.75), there was no statistically significant correlation between the cation/anion molar ratio and [H^+] from either ion balance or model
predictions. Therefore, the molar ratio does not appear to be a suitable tool to infer the acidity of atmospheric particles at the study site, but it could be a good parameter to distinguish between alkaline and acidic particles.

Data on gaseous NH$_3$ were not available, but the potential relationship was also investigated between NH$_3$/NH$_4^+$ phase partitioning (with NH$_3$ values predicted by ISORROPIA-II) and particles pH. The hypothesis of phase partitioning equilibrium is justified by the fact that the sampling time-interval (36 h) was much longer than the equilibration time for submicron particles (seconds to minutes; Meng et al., 1995). As previously mentioned, in the present work the number of samples with pH < 0 was significantly lower compared to those with pH > 0. Moreover, ISORROPIA-II predicted that in the 94.5-612 nm size range there would be a significant NH$_3$ fraction in the gas phase.

The detailed NH$_3$/NH$_4^+$ partitioning as a function of RH is presented below, and considerations on the potential effects on the partitioning brought about by changes in the SO$_4^{2–}$ and NO$_3^–$ concentrations, and by temperature affecting both SO$_4^{2–}$ and NO$_3^–$ production, is given in Section S 8 of the SM. The potential role played by temperature on NH$_3$/NH$_4^+$ partition seems to be minimal, but one should also consider that highly acidic aerosols will affect a variety of processes and definitely the partitioning of HNO$_3$ to the gas phase, producing low nitrate aerosol levels. ISORROPIA-II runs predicted gas-phase NH$_3$ concentrations in Iasi as high as $0.52 \pm 0.28$ (0.46) [mean ± stdev (median)] µg m$^{-3}$ at RH < 40 %, $0.61 \pm 0.26$ (0.49) µg m$^{-3}$ at RH = 40–60 %, and $0.96 \pm 0.54$ (0.92) µg m$^{-3}$ at RH > 60 %. These warm-season values are smaller than those reported in a modelling study by Backes et al. (2016), who predicted NH$_3$ abundances as high as 1.6 to 2.4 µg m$^{-3}$ (data extracted from NH$_3$ concentration for the reference case, i.e., FIG. 3 in Backes et al., 2016, for north-eastern Romania). In contrast, the 0.96 ± 0.54 (0.92) µg m$^{-3}$ value at RH > 60 %, which would be mainly found during the cold seasons, seems to be in reasonable agreement with the ≤ 0.8 µg m$^{-3}$ value modelled by Backes et al. (2016) over winter-time. Moreover, from ISORROPIA-II runs performed at RH < 40 %, it was estimated that (77.6 ± 28.4) % or (79.3 ± 26.2) % (mean ± stdev) of the NH$_3$ predicted by ISORROPIA-II could be present in the gaseous phase (with reference to both NH$_4^+$ derived from raw IC data, and to the NH$_4^+$ (total) fraction). Similar but slightly decreasing values were predicted for gas-phase NH$_3$ as RH increased, namely (76.5 ± 30.9) % or (78.6 ± 28.2) % (raw IC and NH$_4^+$ (total), respectively) at RH = 40–60 % and (68.3 ± 36.7) % or (74.5 ± 29.7) % at RH > 60 %.

In other studies, thermodynamic equilibrium calculations predicted that all of the NH$_3$ was mainly susceptible of partitioning to the particle phase at the equilibrium, and also that > 44 or 51 % of the investigated samples presented aerosols pH < 0 (Hennigan et al., 2015). However, it seems that at the investigated Romanian site the atmosphere could be rich enough in NH$_3$ so as to allow its occurrence on the gas phase while also promoting particle-phase partitioning. The seasonal trends in the NH$_3$ concentrations derived from ISORROPIA runs for Iasi are reported in FIG. S 1 (Section S 9). The same section reports considerations on possible interrelated emission factors governing the distribution in the NH$_3$ concentration levels in Iasi.

The ISORROPIA-II data referred to the 155–612 nm size range (regardless of RH) suggested that the aerosol ammonium fraction (NH$_4^+/(\text{NH}_3 + \text{NH}_4^+$)) was over 0.20 irrespective of the calculation procedure (raw IC or NH$_4^+$ (total)). Clear (NH$_4^+/(\text{NH}_3 + \text{NH}_4^+$)) maxima were observed at 381 nm (raw IC data: 0.71 at RH < 40 %, 0.63 at RH = 40–60 %, and 0.76
at RH > 60 %; NH$_4^+$ (total), defined as the sum between that derived from raw IC data and the part estimated by using the rationale of Arsene et al. (2011): 0.66 at RH < 40 %, 0.56 at RH = 40–60 %, 0.65 at RH > 60 %). As seen in FIGS. 5c,d, the aerosol pH is very low in the 155–612 nm size range, while it often approaches 8 in the 27.6–94.5 and 612–9940 nm size ranges. Although not presented, in the 155–612 nm size range the aerosol ammonium fraction (NH$_4^+$/(NH$_3$ + NH$_4^+$)) approaches 1, while in other two size ranges it is very low or close to 0 thereby suggesting the occurrence of gaseous NH$_3$.

3.2.2 Seasonality of the major water-soluble ions

Table 3 shows monthly-based statistics for meteorological variables and mass concentrations of PM$_{10}$, PM$_{2.5}$ and major water-soluble ions in PM$_{2.5}$. Compared to the PM$_{2.5}$ fraction, in the PM$_{10}$ fraction we observed increases in the mass concentrations of the following ions (notation for the % increase: min–max (mean)): 13–80 % (35 %) for Cl$^-$; 1–107 % (32 %) for NO$_3^-$; 1–170 % (63 %) for SO$_4^{2-}$; 1–171 % (22 %) for acetate; 4–136 % (22 %) for formate; 0–294 % (27 %) for oxalate; 16–48 % (32 %) for Na$^+$; 6–58 % (20 %) for K$^+$; 1–105 % (12 %) for NH$_4^+$ (total); 28–83 % (46 %) for Mg$^{2+}$; and 33–123 % (61 %) for Ca$^{2+}$. However, the PM$_{10}$ and PM$_{2.5}$ mass concentrations fractions show statistically significant correlation with a ratio of 1.1 (Pearson coefficient of 0.99, p < 0.001). Higher mass concentrations of specific water-soluble ions (Cl$^-$, NO$_3^-$, K$^+$, NH$_4^+$ and, to some extent, SO$_4^{2-}$) were observed during the cold compared to the warm seasons, probably because of the combination of increased strength of pollution sources and meteorological effects (inducing lower mixing heights or even temperature inversion), or due to different chemical/photochemical processing. Although lowering mixing heights over the cold seasons might increase pollutant concentration in the atmosphere, for some species additional phenomena should be taken into account in order to explain their distribution. For particulate SO$_4^{2-}$, high concentrations can be observed during winter and autumn but also in summer, and in the latter case they can be due to higher temperature and solar radiation that enhance photochemical reactions and the atmospheric oxidation potential, because of the elevated occurrence of oxidant species such as ozone, hydroxyl and nitrate radicals. These conditions favour the oxidation of SO$_2$ to particulate SO$_4^{2-}$. Also particulate C$_2$O$_4^{2-}$ was maximum in summer, possibly due to enhanced photochemical processing. Moreover, the maxima observed for SO$_4^{2-}$ during the cold seasons might be due to the intensification of coal burning for heating purposes. Higher abundances of particulate NO$_3^-$, SO$_4^{2-}$, NH$_4^+$ and K$^+$ in winter compared to summer are reported for other European (Schwarz et al., 2012; Voutsa et al., 2014) and non-European sites as well (Sharma et al., 2007). Sharma et al. (2007) also suggest a potential role of CaCO$_3$ in controlling particulate NO$_3^-$ abundance in Kanpur, India. Seasonal variations for selected water-soluble ionic components in PM$_{2.5}$ are shown in FIGS. 6a-h, while FIG. 6i reports the variation of the mixing layer depth at the investigated site. Fine particulate Cl$^-$, NO$_3^-$, K$^+$, NH$_4^+$ (total), and to some extent even SO$_4^{2-}$, seem to exhibit distinct seasonal variations with maxima during the cold and minima over the warm seasons, which might be related to changes in the mixing layer depth. The summer minima observed for both NO$_3^-$ and NH$_4^+$ are not surprising, because NH$_4$NO$_3$, is volatile and tends to dissociate to gas-phase NH$_3$ and HNO$_3$ at high temperatures. Coarse particulate C$_2$O$_4^{2-}$, Ca$^{2+}$ and Na$^+$ did not show much variation with respect to seasons. However, SO$_4^{2-}$ and C$_2$O$_4^{2-}$ showed similar patterns (implying most probably common sources), and the Ca$^{2+}$ trend suggests prevalent contribution from soil
dust. Higher ion concentrations in winter than in summer are reported by Sharma et al. (2007) for Kanpur (India), while Ianniello et al. (2011) report opposite trends for Beijing (China).

Particulate Cl– mass concentrations show a clear seasonal pattern, with higher values during the cold than during the warm seasons (FIG. 6a). The chloride mass concentration in both PM<sub>2.5</sub> and PM<sub>10</sub> had a statistically significant correlation with RH, temperature (only for PM<sub>10</sub> fraction), particle loading and mixing layer depth (detailed statistics in Table S 4). The chloride maxima during the cold seasons might be the result of increased coal burning for heating purposes or of the use of NaCl in winter on icy/snowy roads. These observations are in agreement with other studies in eastern European sites (Arsene et al., 2011; Alastuey et al., 2016). However, the Cl– mass concentration follows a similar pattern as that of K+ (tracer of biomass burning), thereby suggesting that that over the cold seasons wood burning might become an important heating source (Christian et al., 2010; Akagi et al., 2011).

Also nitrate shows cold-season maxima and warm-season minima (see Table 3 and FIG. 6b). The inset distribution presented within NO<sub>3</sub>– seasonal variation (FIG. 6b) suggests that, in summer, the coarse PM fraction can bring significant contributions to the aerosol atmospheric burden of nitrate. In our work, fine particulate NO<sub>3</sub>– mass concentrations varied from 0.31 to 3.62 µg m<sup>–3</sup> (Table 3) and these data are in very good agreement with those predicted for Europe in a modelling study performed by Backes et al. (2016). The data obtained in the present work over the cold seasons (3.62 ± 1.10 µg m<sup>–3</sup> in January, February, December as the coldest months of the year, and 2.65 ± 0.38 µg m<sup>–3</sup> over January, February, March, October, November, and December) seem to be in reasonably good agreement with those predicted for Europe by Backes et al. (2016). In contrast, the 0.59 ± 0.30 µg m<sup>–3</sup> NO<sub>3</sub>– concentration in PM<sub>2.5</sub> measured over the warm seasons (including the April month characterised by predominant air-mass buoyancy, and decreasing down to 0.44 ± 0.12 µg m<sup>–3</sup> if the April month is excluded) is lower than that predicted by Backes et al. (2016) (abundances as high as 0.8 µg m<sup>–3</sup> over summer). Similarly to the NH<sub>4</sub>+ case, this might reflect the susceptibility of NO<sub>3</sub>– to be transferred to the gas phase over the warm seasons. The NO<sub>3</sub>– mass concentrations in both PM<sub>2.5</sub> and PM<sub>10</sub> had a statistically significant correlation with RH, temperature, mixing layer depth and particles loading (Table 3, detailed statistics in Table S 4). However, it has to be observed that highly acidic aerosols expected over all seasons have the potential to affect the partitioning of HNO<sub>3</sub> to the gas phase, producing low nitrate aerosol levels. Guo et al. (2015) also report low nitrate aerosol levels during summer. Moreover, NO<sub>3</sub>– heterogeneous formation (i.e., condensation or absorption of NO<sub>2</sub> in moist aerosols or N<sub>2</sub>O<sub>5</sub> oxidation and HNO<sub>3</sub> condensation) generally relates to RH and the particulate loading (Wang et al., 2006; Ianniello et al., 2011). At the investigated site, this process might be of similar importance as the gas–particle conversion, which implies oxidation of precursor gases, such as NO<sub>x</sub>, to nitrate via HNO<sub>3</sub> formation and involving photochemical processes. The high concentration of NO<sub>3</sub>– during the cold seasons might also be caused by higher NH<sub>3</sub> atmospheric levels from yet unaccounted sources, which could neutralize gas-phase H<sub>2</sub>SO<sub>4</sub> and HNO<sub>3</sub> to produce ammonium salts (vide infra). Reactive nitrogen species are emitted to the atmosphere mainly in the forms of NO<sub>x</sub> (from transport or power generation) and NH<sub>3</sub> (agriculture). In Iasi, the animal husbandry sector (open and closed barns, manure storage/spreading) is most likely an important NH<sub>3</sub> source. Moreover, the high relative humidity observed in the cold seasons could offer suitable conditions for significant fractions of
HNO3 and NH3 to be dissolved in humid particles, therefore enhancing particulate-phase NO3\(^-\) and NH4\(^+\) (Pathak et al., 2009, 2011; Ianniello et al., 2010; Sun et al., 2010). From measurements performed in October 2004 in Beijing, China, Kai et al. (2007) also concluded that high RH, stable atmosphere and high NH3 levels can enhance transformation of NOx into NO3\(^-\). Particulate SO4\(^{2-}\) maxima are observable during both cold and warm seasons. However, the particulate SO4\(^{2-}\) mass concentrations showed statistically significant correlation with the measured meteorological parameters only at a 68% confidence level (detailed statistics in Table S 4). The data presented in FIG. 6c show that the seasonal trend of the monthly SO4\(^{2-}\) mean mass concentrations is not as clear as that of NO3\(^-\) and NH4\(^+\), which might suggest the occurrence of regional SO4\(^{2-}\) sources as well (Wang et al., 2016). Moreover, high RH (in Iasi, especially during cold months) may aid the conversion of SO2 to SO4\(^{2-}\) (Kadowaki, 1986), with a significant enhancement of SO4\(^{2-}\) production rate in aqueous phase (Sharma et al., 2007). However, the oxidation of SO2 to sulfate may be induced not only by H2O2 in the aqueous phase, but also by gas-phase radical hydroxyl (Vione et al., 2003). This issue can explain the PM sulfate maxima during the warm seasons because of higher sunlight irradiance and temperature (Stelson and Seinfeld, 1982; Stockwell and Calvert, 1983; Kadowaki, 1986; Wang et al., 2005).

The data reported in Table 3 and FIG. 6f show that particulate NH4\(^+\)(total) has a clear seasonal pattern with maxima during the cold and minima over the warm seasons, in agreement with reports at other European (Schwarz et al., 2012; Bressi et al., 2013; Tositti et al., 2014; Voutsa et al., 2014) or non-European sites (Sharma et al., 2007; Wang et al., 2016). This behaviour is opposite to that reported by Ianniello et al. (2011). The cold-season maxima and warm-season minima we observed can be due to the effect of variations in the mixing layer depth, combined with gas-phase transfer of NH4NO3 to NH3 and HNO3 as temperature increases. In our dataset, NH4\(^+\)(total) varied from 0.8 to 2.09 µg m\(^{-3}\), a much lower range than that reported by Meng et al. (2011) for a more polluted site (Beijing, China, with concentrations varying between 4.73 to 9.04 µg m\(^{-3}\) among various seasons). However, the data we measured in the cold season (2.03 ± 0.30 µg m\(^{-3}\) over January, February and December; 1.65 ± 0.23 µg m\(^{-3}\) over January, February, March, October, November and December) seem to be in reasonable agreement with those predicted for Europe by Backes et al. (2016). In contrast, the 0.90 ± 0.09 µg m\(^{-3}\) NH4\(^+\)(total) measured concentration over the warm seasons is much lower than that predicted by Backes et al. (2016), and the discrepancy might actually reflect the limitation of the experimental measurement techniques concerning NH4NO3 volatility. Moreover, the NH4\(^+\)(total) mass concentration correlated significantly with RH and particle loading in both the PM\(_{2.5}\) and PM\(_{10}\) fractions, and it anticorrelated significantly with temperature and mixing layer depth (detailed statistics in Table S 4). The PM\(_{2.5}\) fraction showed statistically significant correlation also with the mixing depth (Pearson coefficient higher than 0.67, p = 0.016). The seasonal variation of particulate NH4\(^+\)(total) follows especially those of particulate NO3\(^-\) and Cl\(^-\), which would indicate that most probably NH4\(^+\)(total) largely originates from neutralization between NH3, HNO3 and HCl (Wang et al., 2006), or that the cited particulate species derive from similar gas-to-particle processes (Huang et al., 2010). Although in the present work gaseous NH3 was not measured, ISORROPIA-II runs predicted that the atmosphere was often in a gaseous ammonia-rich state, independently of the RH values (\([\text{NH}_3]/([\text{HNO}_3] + [\text{HCl}]) \gg 1\)).
At the investigated site, C$_2$O$_4^{2-}$ and SO$_4^{2-}$ show similar behaviour and the C$_2$O$_4^{2-}$ maxima during summer may suggest photochemical and/or biogenic contributions to its abundance (Laongsri and Harrison, 2013). The Na$^+$ ion, tracer of sea-salt or NaCl aerosols, shows higher concentrations during spring when one has a predominant long-range transport of air masses from the S-SE sector, with contributions from natural sources and especially sea-spray aerosols from the Black Sea. Particulate K$^+$ mass concentrations also show a clear seasonal pattern, with higher values during the cold compared to the warm seasons (FIG. 6e). This phenomenon could be due to increased wood burning combined with a limited mixing layer depth. However, particulate K$^+$ mass concentrations also had some maxima when one expects intense agricultural biomass burning for field clearing (i.e., April, July, and September). The K$^+$ mass concentrations follow a similar pattern as Cl$^-$ (Pearson coefficient of 0.79, $p = 0.002$) but a different one compared to SO$_4^{2-}$. Therefore, we suggest that intense wood burning may be a common source for K$^+$ and Cl$^-$ species in the study area (Christian et al., 2010; Akagi et al., 2011).

High mass concentrations of Ca$^{2+}$ and Mg$^{2+}$ (with Mg$^{2+}$ shown in Table 3 but not in FIG. 6), as soil/dust tracers, were observed especially during spring and summer. Over these seasons, lower precipitation frequency and high wind speed contribute to the observed behaviour. During the cold seasons, low wind speeds might prevent mineral dust resuspension and produce low values for these ions. However, Mg$^{2+}$ and Ca$^{2+}$ as mineral ions did not correlate with either PM$_{2.5}$ or PM$_{10}$, which suggests that inorganic particles would be mainly produced by NH$_3$-triggered secondary processes.

### 3.2.3 Stoichiometry of (NH$_4$)$_2$SO$_4$, NH$_4$NO$_3$ and NH$_4$Cl

Table 4 presents the correlation matrix (Pearson coefficients) for the major ionic species (Cl$^-$, NO$_3^-$, SO$_4^{2-}$, CH$_3$COO$^-$, HCOO$^-$, C$_2$O$_4^{2-}$, HCO$_3^-$, Na$^+$, NH$_4^+$ (total), K$^+$, Mg$^{2+}$, Ca$^{2+}$) in PM$_{2.5}$, for both the cold (Table 4.a) and the warm seasons (Table 4.b). Despite similar correlations in PM$_{10}$ as well, PM$_{2.5}$ was selected for the correlation matrix analysis because of higher representativeness. For the cold seasons there are significant correlations (at the 99.9 % confidence level) among many chemical pairs, suggesting an important occurrence of (NH$_4$)$_2$SO$_4$, NH$_4$NO$_3$ and (NH$_4$)$_2$C$_2$O$_4$. In the warm seasons, (NH$_4$)$_2$SO$_4$, Mg(NO$_3$)$_2$, and NaCl seem to be the most important. However, Ca(HCO$_3$)$_2$ might play a role during both cold and warm seasons. Over the cold seasons, K$^+$ shows statistically significant correlation with many inorganic (NO$_3^-$, Cl$^-$ and SO$_4^{2-}$) and organic (HCOO$^-$ and C$_2$O$_4^{2-}$) anions, thereby suggesting that at least some of these species might have common biomass burning sources (Ianniello et al., 2011 and references therein). However, in the case of SO$_4^{2-}$ and NO$_3^-$ the most likely explanation would rather be the use of both coal and wood for burning, as well as the effect of the mixing layer depth. In the ambient atmosphere, inorganic ammonium salts such as NH$_4$HSO$_4$, (NH$_4$)$_2$SO$_4$, NH$_4$NO$_3$ and NH$_4$Cl are known to be produced by gas-to-particle conversion processes. In the present work, from the ionic balance analysis the NH$_4^+$ ion was assigned as the most critical parameter in the chemical composition analysis of aerosol particles (with 274 analysed samples, i.e., about one quarter of the total, being highly deficient in cations). FIGURES 7a,b present the relationship between the molar concentrations of fine particulate NH$_4^+$ (i.e., from raw IC data and also in the total form estimated under the assumptions from Arsene et al., 2011) and that of particulate SO$_4^{2-}$ for both the cold (FIG. 7a) and warm (FIG. 7b) seasons. Correlations between particulate NH$_4^+$ and SO$_4^{2-}$ are statistically significant in both cases (detailed statistics in Table S 5).
During cold and warm seasons, the \([\text{NH}_4^+]/(2\times[\text{SO}_4^{2-}])\) molar ratio was either ~1 (raw IC data), or (NH\(_4^+\) (total) values) equal to 1.76 (cold seasons) and 1.02 (warm seasons). These data suggest the existence of enough NH\(_3\) for the complete neutralization of H\(_2\)SO\(_4\), and also a predominance of particulate (NH\(_4\))\(_2\)SO\(_4\) in agreement with the observations of Ianniello et al. (2011). Moreover, as shown in Table 4, for particulate SO\(_4^{2-}\) and NH\(_4^+\) (total) the correlation was statistically significant (with Pearson coefficients of 0.96, \(p < 0.001\) for cold and 0.97, \(p < 0.001\) for warm seasons), thereby suggesting that (NH\(_4\))\(_2\)SO\(_4\) could be formed from H\(_2\)SO\(_4\)(g) and NH\(_3\)(g) in either case. However, the 1.76 value for the \([\text{NH}_4^+]\) (total)/(2×[SO\(_4^{2-}\)]) molar ratio during the cold seasons will indicate that there should still be particulate NH\(_4^+\) potentially available for combination with other anions (or the existence of enough excess ammonia to neutralize acidic species such as HNO\(_3\) and HCl). Zhao et al. (2016) report a \([\text{NH}_4^+]\)/(2×[SO\(_4^{2-}\)]) molar ratio of 1.54 (\(R^2 = 0.63\)), indicating the complete neutralization of H\(_2\)SO\(_4\) and a predominance of (NH\(_4\))\(_2\)SO\(_4\) in sulfate salts during the cold seasons.

Unfortunately, at present, no measured NH\(_3\) values are available for the interest site but it is still believed that, in the atmosphere of Iasi, sufficient gas-phase NH\(_3\) occurs to promote both the homogeneous and heterogeneous formation of nitrate salts in the collected aerosol particles. The NH\(_4\)NO\(_3\) formation routes might involve either the homogeneous reaction between gaseous HNO\(_3\) and NH\(_3\) (Ianniello et al., 2011), or the heterogeneous reaction between NH\(_3\) and the products formed upon hydrolysis of N\(_2\)O\(_5\) that could be present on the surface of the pre-existing moist aerosols under relatively high humidity (Pathak et al., 2011; Shon et al., 2013). Actually, gaseous NH\(_3\) can influence both the inorganic ions and the aqueous-phase H\(^+\) distribution in aerosols. The concentration of H\(^+\) in aqueous aerosols is mainly determined by the balance of the acidic ionic components with the basic ones. During both cold and warm seasons, about half or slightly more than half of the collected samples were found alkaline with pH values fluctuating between 7 and 8. The remaining samples were acidic and with pH values ranging mainly from 1 to 3. Zhao et al. (2016) report that, on average, a +25 % perturbation in the NH\(_3\) level could lead to a 0.14 unit pH increase, and a – 25 % perturbation could cause a 0.23 unit pH decrease. They concluded that sufficient NH\(_3\) was frequently present in wintertime atmosphere, and also that the fine collected particulates were almost fully neutralized by NH\(_3\).

FIGURES 7c,d,e,f show the relationships between the fine-particulate molar concentrations of (i) NH\(_4^+\) and the sum of SO\(_4^{2-}\) and NO\(_3^-\) (FIGS. 7c,d), and (ii) NH\(_4^+\) and the sum of SO\(_4^{2-}\), NO\(_3^-\) and Cl\(^-\) (FIGS. 7e,f), for both cold and warm seasons. From details given in FIGS. 7c,d,e,f it can be easily observed that NH\(_4^+\), with both calculation methods (IC and total), was in deficit over the cold and warm seasons. Under these circumstances, it is believed that both particulate NO\(_3^-\) and Cl\(^-\) could be associated with other alkaline species or be part of acidic aerosol. Because the neutralising capacity of NH\(_4^+\) toward SO\(_4^{2-}\), NO\(_3^-\) and Cl\(^-\) acidic species might give a rough indication on the potential particles acidity (Li et al., 2015), from FIGS. 7c,d,e,f it is quite clear that at the study site, if NH\(_4^+\) derived from raw IC data is used, the neutralisation ratios in the investigated particles are less than unity. This suggests that the atmospheric particles are most likely acidic, and also that a more complex chemistry is ongoing involving the HNO\(_3\) and HCl species. However, when NH\(_4^+\) (total) is used, the neutralisation ratio approaches 1 and suggests a possibly complete neutralization of particles acidity. From details in FIGS. 7e,f it can be easily seen that actually the available NH\(_4^+\) is not enough to compensate for other species. However, it should
be noted that Cl$^-$ is not significantly influencing the neutralization of particles acidity. In a study performed by Zhao et al. (2016) the authors reported a $[\text{NH}_4^+] / (2 \times [\text{SO}_4^{2-}] + [\text{NO}_3^-])$ molar ratio of 0.86 ($R^2 = 0.78$) and an $[\text{NH}_4^+] / (2 \times [\text{SO}_4^{2-}] + [\text{NO}_3^-] + [\text{Cl}^-])$ molar ratio of 0.60 ($R^2 = 0.86$). Details presented in FIGS. 7c,d clearly show that when NH$_4^+$ (total) is taken into account, a complete neutralization of H$_2$SO$_4$ and HNO$_3$ can be achieved during the cold seasons (FIG. 7c). In contrast, during the warm seasons (FIG. 7d) the molar ratio is slightly lower than 1 (i.e., 0.95). According to Seinfeld and Pandis (1998), during the warm seasons the high temperature and the low relative humidity would be favourable for NH$_4^+$ to reach a minimum concentration, because it is mainly transformed into NH$_3$. Actually, temperature values above 25 °C, such as those often encountered at the investigated site over the warm seasons, are known to prevent formation of particulate NH$_4$NO$_3$ (Adams et al., 1999). Under these circumstances, a considerable decrease in the $[\text{NH}_4^+]$(total)/($[\text{NO}_3^-] + 2 \times [\text{SO}_4^{2-}]$) molar ratio is to be expected during warm seasons. The cold-season temperature and relative humidity at the investigated site were, respectively, 5.3 ± 3.9 °C and (65.3 ± 12.8) %. Coherently, the data presented in Table 4 show that the ($[\text{NO}_3^-], \text{NH}_4^+$(total)) pair has significant correlation (Pearson coefficient of 0.98, $p < 0.001$) only during the cold seasons, while during the warm seasons (temperature and relative humidity of, respectively 18.9 ± 3.8 °C and 40.5 ± 7.7 %) the correlation is very poor as increasing temperature and decreasing relative humidity limit the production of the NH$_3$NO$_3$ aerosol (Matsumoto and Tanaka 1996; Utsunomiya and Wakamatsu 1996; Alastuey et al., 2004).

In the atmosphere, both H$_2$SO$_4$ and HNO$_3$ are known to compete for the reaction with NH$_3$ to form (NH$_4$)$_2$SO$_4$ and NH$_4$NO$_3$. As presented in Sect. 3.2.2, at the investigated site a reduction in both in NO$_3^-$ and SO$_4^{2-}$ was observed over the warm seasons and this may cause a further increase in gas-phase NH$_3$. It is also interesting to observe that the reaction rate constant of (NH$_4$)$_2$SO$_4$ aerosol formation is similar to the rate constant of NH$_4$NO$_3$ formation (Harrison and Kitto, 1992; Pandolfi et al., 2012; Behera et al., 2013), and both are much higher than the rate constant between NH$_3$ and HCl (Behera and Sharma, 2012). This issue probably dictates the competition of any of the particulate SO$_4^{2-}$, NO$_3^-$ and Cl$^-$ for the available NH$_4^+$. Usually, when a sufficient amount of NH$_3$ is available for neutralization of H$_2$SO$_4$ and HNO$_3$, fine-mode (NH$_4$)$_2$SO$_4$ and NH$_4$NO$_3$ will be formed via reactions R1 (Cziczo et al., 1997; Zhang et al., 2015) and R2 (Fountoukis and Nenes, 2007; Zhang et al., 2015),

\[
2\text{NH}_3(g) + \text{H}_2\text{SO}_4(aq) \rightleftharpoons (\text{NH}_4)_2\text{SO}_4(aq) \quad (R1)
\]

\[
\text{NH}_3(g) + \text{HNO}_3(g) \rightleftharpoons \text{NH}_4\text{NO}_3(s) \quad (R2)
\]

In contrast, in NH$_3$-limited environments, coarse-mode NO$_3^-$ is formed through reaction R3 involving Mg$^{2+}$ (but not Ca$^{2+}$):

\[
\text{MgCO}_3(aq) + 2\text{HNO}_3(g) \rightleftharpoons \text{Mg(NO}_3)_2(s) + \text{H}_2\text{O} + \text{CO}_2(g) \quad (R3)
\]

During the warm seasons, however, higher concentrations of (NH$_4$)$_2$SO$_4$ compared to NH$_4$NO$_3$ are expected because (NH$_4$)$_2$SO$_4$ is less volatile than NH$_4$NO$_3$ (Utsunomiya and Wakamatsu, 1996). Moreover, NH$_4$NO$_3$ will be formed only when excess NH$_3$ is available to react with HNO$_3$. A modelling study by Backes et al. (2016) suggests that a reduction of NH$_3$ emissions by 50 % may lead to a 24 % reduction of the total PM$_{2.5}$ concentrations in northwest Europe, mainly due to reduced formation of NH$_4$NO$_3$. However, the NH$_3$ concentration in the atmosphere over Europe seems to be high enough to saturate the reaction forming (NH$_4$)$_2$SO$_4$ particles, even in a scenario of reduced NH$_3$ levels, while on the contrary it is not
high enough to saturate the reaction with HNO₃ to form NH₄NO₃ particles. A reduced formation of NH₄NO₃ particles may lead to an increase in gas-phase HNO₃ during winter. In our study, results from ISORROPIA-II thermodynamic model foresee an increase of gas-phase HNO₃ at higher RH values during cold seasons. Higher levels of gas-phase HNO₃ may increase its condensation onto existing particles such as sodium chloride (NaCl), and the replacement of Cl⁻ with NO₃⁻ may enhance the concentration of HCl in the atmosphere (similar processes are described in Arsene et al., 2011).

In the atmosphere, additional non-volatile species containing nitrate and chloride might also be present, thus we investigated the potential of fine-particulate NO₃⁻ and Cl⁻ to be chemically bound to Ca²⁺, Mg²⁺, K⁺ or Na⁺. The free NO₃⁻ and Cl⁻ concentrations, defined as the fractions of excess nitrate and chloride that are not bound to alkali or alkaline earth metals, were estimated for both the cold and warm seasons according to the concept described in Ianniello et al. (2011). Zero or negative values of free NO₃⁻ and Cl⁻ imply that NH₄NO₃ and NH₄Cl are not present. Estimated free NO₃⁻ and Cl⁻ concentrations showed similar contributions in both the PM2.5 and PM10 fractions. Over the cold seasons we calculated \(6.5 \pm 7.9 \times 10^{-3} \mu \text{mol m}^{-3} (0.4 \pm 0.5 \mu \text{g m}^{-3})\) for NO₃⁻, and negative values for free Cl⁻; over the warm seasons we had \(1.6 \pm 1.8 \times 10^{-3} \mu \text{mol m}^{-3} (0.1 \pm 0.1 \mu \text{g m}^{-3})\) for NO₃⁻, and again negative values for free Cl⁻ (data provided as mean ± stdev). This result suggests the potential presence of NH₄NO₃, especially during the cold seasons, but it excludes the occurrence of NH₄Cl. During the cold seasons, particulate NO₃⁻ didn’t show correlation with either Ca²⁺ or Mg²⁺, but it showed significant correlation with K⁺ (\(r = 0.85, p < 0.001\), see Table 4), which indicates the possible formation of the non-volatile KNO₃ salt along with NH₄NO₃. Over the warm seasons, fine particulate NO₃⁻ didn’t show correlation with K⁺, but it showed significant correlation with Na⁺ and Mg²⁺ (respectively, \(r = 0.71, p < 0.001\) and \(r = 0.86, p < 0.001\)), indicating possible formation of non-volatile NaNO₃ and Mg(NO₃)₂ but not of Ca(NO₃)₂ salts. Moreover, in the warm seasons the fine-particulate NO₃⁻ also showed statistically significant correlation with HCO₃⁻ (\(r = 63, p < 0.001\)), which suggests a prevalence of particulate NO₃⁻ formation via the mineral route over the homogeneous reactions. The interaction between NO₃⁻ and Mg²⁺ could increase in importance during the warm seasons because, as NH₄⁺ is not available, neutralization of HNO₃ could occur on coarse soil particles rich in Mg²⁺ (Matsumoto and Tanaka, 1996; Utsunomiya and Wakamatsu, 1996; Alastuey et al., 2004).

From the literature it is known that species such as (NH₄)₂SO₄ and NH₄HSO₄, at room temperature, uptake water (deliquesce) at RH values of, respectively, \((79 \pm 1)\%\) and \(39\%\) (Cziezo et al., 1997). The (NH₄)₂SO₄ aerosol particles might remain in “metastable” or supersaturated state liquid phase until a very low RH (crystallization point, RH around 33%) is reached (only thereafter a solid might be formed). In contrast, for NH₄HSO₄ it has been shown that the solid phase is difficult to be formed. In the present work, it is very likely that only over July (RH of 32.95 %), August (RH of 37.12 %) and September (RH of 31.56 %) the formation of solid (NH₄)₂SO₄ or NH₄HSO₄ could occur.

Pure NH₄NO₃ deliquesces at 62 % RH and there is suggestion that sometime even at 8 % RH the crystallization point is not reached (Dougle et al., 1998). According to suggestions from the literature, only during months when the ambient RH is lower than the relative humidity at deliquescence (RHD), the NH₄NO₃ is considered as a solid (Seinfeld and Pandis, 1998). In the present work, from March to October the ambient RH was always lower than the RHD and NH₄NO₃ could thus be assumed to be in equilibrium with the solid phase. Within all the other months, from the estimated RHD values there is the
possibility that NH$_4$NO$_3$ is in equilibrium with the aqueous phase and deliquescent particles. However, in the investigated site, solid NH$_4$NO$_3$ could be formed almost all over the year either due to the very complex chemical composition of the collected particles, or due to abundant contribution of organic carbon to the particles mass concentration (Dougle et al., 1998). Formation of NH$_4$NO$_3$ over the warm seasons has been reported also by Ianniello et al. (2011) but under different conditions. For deliquesced particles there is suggestion that most of the fine particulate NO$_3^-$ occurs as an internal mixture with SO$_4^{2-}$, and also that HNO$_3$ can easily be absorbed into the droplets (Huang et al., 2010). In specific circumstances, the fine particulate NO$_3^-$ can be formed from HNO$_3$ and NH$_3$ through heterogeneous reactions on fully neutralized fine particulate SO$_4^{2-}$, which is abundant in urban areas (Stockwell et al., 2000). In the present work, a statistically significant correlation between SO$_4^{2-}$ and NO$_3^-$ was observed during the cold seasons in the PM$_{2.5}$ fraction (r = 0.91, p < 0.001). High concentrations of NO$_3^-$ were found in the presence of elevated RH levels (significant correlation: r = 0.84, p < 0.001), while SO$_4^{2-}$ concentrations were high over the entire RH range (non-significant correlation: r = 0.44, p = 0.177). These results can be interpreted as nitrate being produced on pre-existing sulfate aerosols, which could provide sufficient surface area and water content for the heterogeneous reactions to occur. Although the formation of fine particulate NO$_3^-$ can take place via reaction (R2), in particular circumstances and especially at high RH values, the amounts of the gaseous precursors such as NH$_3$ and HNO$_3$ may have relatively little influence on the fine-particulate NO$_3^-$ formation (Markovic et al., 2011).

The salt NH$_4$Cl is known to be 2–3 times more volatile than NH$_4$NO$_3$, as HCl is more volatile than HNO$_3$. Moreover, at RH < 75–85 %, particulate NH$_4$Cl is in equilibrium with the gaseous compounds (Ianniello et al., 2011 and references therein). Coherently, during the cold seasons particulate NH$_4^+$ (total) showed a statistically significant correlation with particulate Cl$^-$ (Pearson coefficient of 0.73, p < 0.001), but during the warm seasons the correlation was very poor. Finally, only during the cold seasons significant correlations were observed between fine particulate Cl$^-$ and SO$_4^{2-}$ (Pearson coefficient of 0.59, p < 0.001) and between fine particulate Cl$^-$ and RH (Pearson coefficient of 0.71, p = 0.010). Usually, high concentrations of fine particulate Cl$^-$ and SO$_4^{2-}$ were found at high levels of RH (35–83 %). Under these circumstances, the amount of gaseous precursors is believed to have relatively little influence on the formation of fine particulate Cl$^-$. If formed, NH$_4$Cl is most probably generated from HCl and NH$_3$ through heterogeneous reactions on neutralized sulfate particles. However, in the present work, estimation of free Cl$^-$ suggests that no Cl$^-$ was available to be bound with other chemical species (i.e., NH$_4^+$) apart from alkali or alkaline earth metals, thereby excluding a significant occurrence of NH$_4$Cl (especially over the warm seasons).

3.3 Relative ionic contribution in size resolved aerosol particles from Iasi and potential influence of long-range transport phenomena on particles size distribution

FIGURES 8a,b,c,d present, as monthly based averages, the relative contributions of identified and quantified water-soluble ions to total detected components in fractions grouped in four stages, i.e., 0.0276–0.0945 μm size range (FIG. 8a), 0.155–0.612 μm size range (FIG. 8b), 0.946–2.39 μm size range (FIG. 8c) and 3.99–9.94 μm size range (FIG. 8d). From details presented in FIG. 8a, for the 0.0276–0.0945 μm fraction there is important contribution of formate, acetate and oxalate that
may actually indicate a possible important role of organic acids in secondary organic aerosols formation. Higher values of these components over the warm seasons may suggest an enhancement in the role of biogenic emission sources. Important contributions are brought by SO$_4^{2-}$, NH$_4^+$ (total), K$^+$ and even by the (unexpectedly high) HCO$_3^-$.

However, high particulate HCO$_3^-$ is also evident for the 0.946–2.39 µm (FIG. 8c) and 3.99–9.94 µm size range (FIG. 8d) fractions. The 0.155–0.612 µm fraction (FIG. 8b) seems to be mainly constituted by SO$_4^{2-}$, NO$_3^-$ and NH$_4^+$ (total), with very small contributions from other ions.

The seasonal variation observed mainly for SO$_4^{2-}$ and NO$_3^-$ might suggest an enhancement of photo-oxidative processes over the warm seasons. The 0.946–2.39 µm (FIG. 8c) and 3.99–9.94 µm size range (FIG. 8d) fractions have a non-homogeneous chemical composition dominated mainly by HCO$_3^-$, NO$_3^-$ and organics. Among all the analysed ions in the investigated period, in the PM$_{10}$ fraction SO$_4^{2-}$ was the most abundant with (26.0 ± 4.3) % contribution, followed by NO$_3^-$ (26.0 ± 10.7 %), NH$_4^+$ (total) (15.0 ± 3.4 %), organics including acetate, formate and oxalate (12.2 ± 3.7 %) and HCO$_3^-$ (10 ± 7.7 %). Similarly, in the PM$_{2.5}$ fraction SO$_4^{2-}$ was again the most abundant (28.9 ± 5.6 %) followed by NO$_3^-$ (19.6 ± 12.1 %), NH$_4^+$ (total) (16.6 ± 3.2 %), organics including acetate, formate and oxalate (11.4 ± 4.0 %) and HCO$_3^-$ (8.0 ± 6.8 %). In both the PM$_{10}$ and the PM$_{2.5}$ fractions, the largest contribution of SO$_4^{2-}$ was observed in June 2016 with, respectively, (34.6 ± 10.9 %) and (40.8 ± 11.0 %).

During the cold seasons, particulate SO$_4^{2-}$ and NO$_3^-$ contributions were, respectively, (23.6 ± 2.3 %) and (28.6 ± 4.9 %) for PM$_{10}$ and (25.5 ± 2.9 %) and (30.1 ± 5.8 %) for PM$_{2.5}$. During the warm seasons, the SO$_4^{2-}$ and NO$_3^-$ contributions, were, respectively, (28.5 ± 4.7 %) and (10.1 ± 4.8 %) for PM$_{10}$, and (32.4 ± 5.6 %) and (9.2 ± 5.4 %) for PM$_{2.5}$. Wonaschutz et al. (2015) report for Vienna (Austria) NO$_3^-$ contributions of 31.3 % during winter and of 6.9 % during summer.

Particulate Ca$^{2+}$ and HCO$_3^-$, as dust tracer ions, brought a significant contribution especially over the warm seasons (43.8 ± 11.2 % in PM$_{10}$ and 37.8 ± 12.3 % in PM$_{2.5}$ in August 2016), suggesting that Ca$^{2+}$ and HCO$_3^-$ mostly originate from soil dust re-suspension during the dry seasons. In 2016, spring was the season with predominant air masses undertaking long-range transport from the S-SE sector (see FIG. 1), thereby suggesting the presence of sea-spray aerosols from the Black Sea or other marine areas. The highest contributions from sea-spray aerosols tracers (i.e., Na$^+$ and Mg$^{2+}$, Masiol et al., 2012) were actually recorded in April (5.8 % for Na$^+$ and 0.7 % for Mg$^{2+}$). Although the most abundant ions are SO$_4^{2-}$, NO$_3^-$ and NH$_4^+$ (total), organics (including acetate, formate and oxalate) might also bring significant contributions ((17.3 ± 4.4) % in PM$_{10}$ and 18.2 ± 5.1 % in PM$_{2.5}$ in July 2016, much higher than that reported by Arsene et al., 2011, for Iasi). The difference might reflect either an inversion of the photochemistry taking place at the investigated location, or differences in the sampling efficiency between the two studies.

FIGURES 9a,b,c,d show the size distributions of seasonal averaged mass concentrations for Cl$^-$, NO$_3^-$, SO$_4^{2-}$, NH$_4^+$ (FIGS. 9a,b) and K$^+$, Na$^+$, Mg$^{2+}$, Ca$^{2+}$ (FIGS. 9c,d). While during the cold seasons NO$_3^-$, SO$_4^{2-}$, NH$_4^+$ and K$^+$ reside mainly in the fine mode with maxima at ~ 381 nm, all the other ions (i.e., Cl$^-$, Na$^+$, Mg$^{2+}$, Ca$^{2+}$) have major contributions in the supermicron mode (maxima between 1.6–2.39 µm). Over the cold seasons, clear evidences were obtained of the occurrence of chloride in a bimodal distribution. During the warm seasons only SO$_4^{2-}$ and K$^+$ presented clear maxima at 381 nm, while all
the other identified/quantified species had more important contributions in the super-micron mode. For \( \text{SO}_4^{2-} \), a larger modal diameter in the cold compared to the warm seasons is most likely due to hygroscopic growth under high RH, and/or to increased secondary aerosol production (lower temperatures facilitate condensation). Secondary aerosol mass from aqueous-phase reactions may also play a role (Wonaschutz et al., 2015). The maxima observed in the coarse mode could also be explained considering that heterogeneous chemistry on dust particles might act as a source for some particulate species (Wang et al., 2012).

While particulate \( \text{NO}_3^- \) over the cold seasons presented monomodal distributions in the sub-micron size range (maxima at 381 nm), over the warm seasons a second mode was observed with maxima in the 1.60-2.39 \( \mu \)m size range. Such a size distribution suggests that \( \text{NO}_3^- \) during the warm seasons is produced by adsorption of \( \text{HNO}_3 \) on sea salt and soil particles (Park et al., 2004). According to Karydis et al. (2016), particulate \( \text{NO}_3^- \) is not associated only with \( \text{NH}_4^+ \) in the fine mode. Light metal ions such as \( \text{Ca}^{2+}, \text{Mg}^{2+}, \text{Na}^+, \text{and K}^+ \), which mainly occur in the coarse mode, can be associated with \( \text{NO}_3^- \) and affect its partitioning into the aerosol phase. Dust effects on the distribution of particulate species might include a decrease of fine-mode \( \text{NH}_4^+ \) and a shift of particulate \( \text{NO}_3^- \) from the fine to the coarse mode (Wang et al., 2012). In addition, the presence of significant levels of \( \text{NO}_3^- , \text{Cl}^- , \text{Mg}^{2+}, \text{Ca}^{2+} \text{and Na}^+ \) in the coarse fraction might suggest that \( \text{NO}_3^- \) possibly originates upon reaction of \( \text{HNO}_3 \) with \( \text{MgCO}_3, \text{CaCO}_3 \text{or NaCl} \). Similar patterns were identified in Vienna, Austria (Wonaschutz et al., 2015) and in Prague, Czech Republic (Schwarz et al., 2012). Significant amounts of particulate \( \text{NO}_3^- \) formed upon reaction of \( \text{HNO}_3 \) with \( \text{CaCO}_3 \) on soil-derived particles have also been observed and reported (Yao et al., 2003; Sharma et al., 2007). Moreover, sea-salt aerosols may also undergo chemical transformation of \( \text{NaCl} \) into \( \text{NaNO}_3 \) during transport (Schwarz et al., 2012).

The size distributions of particulate \( \text{K}^+ \) reflect the occurrence of one dominant fine mode (with maxima at 381 nm) during both the cold and the warm seasons, and of a second less important mode during the warm seasons (with maxima in the 0.946–1.6 \( \mu \)m range). Such behaviour most likely reflects contributions from biomass burning all over the year (Schmidl et al., 2008; Pachon et al., 2013). For both \( \text{Ca}^{2+} \text{and Mg}^{2+} \) ions, clear monomodal mass distributions with maxima in the 1.6 to 2.39 \( \mu \)m size range were observed over the investigated period. Over the warm seasons, \( \text{Ca}^{2+} \) accounts for \( (7.0 \pm 2.9) \% \) of the \( \text{PM}_{10} \) fraction and for \( (5.5 \pm 2.9) \% \) of the \( \text{PM}_{2.5} \) fraction, while over the cold seasons it accounts for only \( (3.0 \pm 0.6) \% \) of the \( \text{PM}_{10} \) and \( (2.2 \pm 0.5) \% \) of the \( \text{PM}_{2.5} \) fraction. These observations indicate that the impact from soil dust re-suspension could be more important during the warm (dry) seasons. Mineral dust may also explain the higher coarse fraction of \( \text{Mg}^{2+} \) (mineral source being \( \text{MgCO}_3 \)).

Clear evidences were obtained in this work that air-mass origin highly influences the aerosol chemical composition at the investigated site. Annual averaged sector contributions, in terms of PM long-range transport, are shown in FIG. S 2 while the seasonal contributions of PM and particulate inorganic/organic ions associated with different air mass origins are reported as Radar charts in FIG. S 3 (within Section S 10). FIG. S 4 (within Section S 10) highlights the % distributions for the identified-quantified ions and the gaseous concentrations of \( \text{NH}_3, \text{HNO}_3 \text{and HCl} \) for selected investigated events, predicted by ISORROPIA-II. Wonaschutz et al. (2015) suggested that in Vienna, Austria, air-mass origin is the most important factor
for bulk PM concentrations, chemical composition of the coarse fraction (> 1.5 µm) and mass size distribution, while it is less important for the chemical composition of the fine fraction (< 1.5 µm). Although Iasi is located far from the Mediterranean or Black Sea, over the warm seasons the sea-salt chloride contribution to the aerosol budget in the area is not entirely excluded (Arsene et al., 2011). Moreover, dust particles originating from the Sahara are acknowledged as travelling across the tropical Atlantic Ocean (10–90 µg m⁻³) and across the Mediterranean, affecting air quality in southern Europe (10–60 µg m⁻³) (Karydis et al., 2016). In the present work, particulate Na⁺ and Cl⁻ ions as tracers of sea-salt aerosols (Tositti et al., 2014) were mainly observed in conditions predominated by contributions from air masses arriving in Iasi from S-SE directions. However, one of the most interesting collected events was that of 9th–11th April 2016. For this event, the PM₁₀ fraction mass concentration was as high as 43.9 µg m⁻³, a value which is about two times higher than the average of the total events. These conditions were highly influenced by air masses originating from both the Saharan desert and the Mediterranean Sea. As shown in FIG. 10a, the size distributions of particulate Na⁺, Ca²⁺, Mg²⁺, Cl⁻ ions and their mass concentrations present a highly dominating mode with maxima at 2.39 µm. For this event, the (Ca²⁺, Mg²⁺) and (Na⁺, Cl⁻) pairs showed statistically significant correlations (respectively, r = 0.94, p < 0.001 and r = 0.85, p < 0.001), suggesting common contributions from mineral Saharan dust (Ca²⁺, Mg²⁺) and from sea-salt marine aerosols (Na⁺, Cl⁻). Moreover, FIG. 10a clearly shows that Na⁺, Ca²⁺, Mg²⁺ and Cl⁻ make a very significant contribution to the total aerosol mass in the super-micron mode, with the maxima at 2.39 µm.

In April 2016, an interesting behaviour was also observed for the averaged mass size distributions of particulate NH₄⁺, NO₃⁻, SO₄²⁻ and Mg²⁺ (FIG. 10b). This month is highly affected by the atmospheric air masses buoyancy phenomenon, as shown by trajectories analysis for selected events and, while particulate NH₄⁺ and SO₄²⁻ were mainly residing in the fine mode with clear maxima at 381 nm, NO₃⁻ and Mg²⁺ also presented a predominant mode in the 1.6–2.39 µm fraction. Such distributions, corroborated with meteorological conditions, would actually suggest a possible heterogeneous formation route for SO₄²⁻ (Wang et al., 2012). In contrast, the adsorption of HNO₃ on mineral dust and sea-salt particles (Karydis et al., 2016) would become more important for NO₃⁻.

4 Conclusions

The atmospheric concentrations of particulate species including acetate, formate, fluoride, chloride, nitrite, nitrate, phosphate, sulfate, oxalate, sodium, potassium, ammonium, magnesium, and calcium were measured over 2016 at an urban site in Iasi, north-eastern Romania. The measurements were carried out by means of a cascade Dekati Low-Pressure Impactor (DLPI), performing aerosol size classification in 13 specific fractions evenly distributed over the 0.0276-9.94 µm size range.

The entire data set was analysed to investigate the seasonal variations in fine particulate species and the meteorological effects, and to examine the contributions of local and regional sources. ISORROPIA-II thermodynamic model runs were
used to estimate the pH values of the collected atmospheric particles, as on the present data-base it was the best method to analyse particles acidity.

Within the aerosol mass concentration, the identified ions mass brings contributions as high as 40.6 % with the rest being unaccounted yet. Fine particulate Cl\(^-\), NO\(_3^-\), NH\(_4^+\) and K\(^+\) exhibited clear seasonal variations with minima during the warm seasons, mainly due to cold-season enhancement in the emission sources, changes in the mixing layer depth and specific meteorological conditions (e.g., higher RH values prevailing in Iasi during the cold seasons). Fine particulate SO\(_4^{2-}\) did not show much variation with respect to seasons. The measured concentrations of NH\(_4^+\) and NO\(_3^-\) in fine-mode (PM\(_{2.5}\)) aerosols were in reasonably good agreement with modelled values for the cold seasons but not for the warm seasons. This observation reflects the susceptibility of NH\(_4\)NO\(_3\) aerosols to be lost due to volatility.

Clear evidences were obtained that NH\(_4^+\) in PM\(_{2.5}\) was primarily associated with SO\(_4^{2-}\) and NO\(_3^-\). However, indirect ISORROPIA-II estimations showed that the atmosphere of Iasi might be ammonia-rich during both cold and warm seasons, so that enough NH\(_3\) would be present to neutralise the H\(_2\)SO\(_4\), HNO\(_3\) and HCl acidic components and to generate fine particulate ammonium salts, in the form of (NH\(_4\))\(_2\)SO\(_4\), NH\(_4\)NO\(_3\) and NH\(_4\)Cl. Significant amounts of fine-particulate NO\(_3^-\) were in fact detected during the cold seasons. The presence of possibly large amounts of NH\(_3\), the domination of (NH\(_4\))\(_2\)SO\(_4\) over NH\(_4\)NO\(_3\) and NH\(_4\)Cl, and the high relative humidity conditions in the cold seasons (likely leading to dissolution of a significant fraction of atmospheric HNO\(_3\) and NH\(_3\)) are among the most important driving forces enhancing the fine-particulate NO\(_3^-\) and NH\(_4^+\) distribution in the investigated site.

Most probably, gaseous NH\(_3\) is not only a precursor of NH\(_4^+\) formation, but it also affects the occurrence of NO\(_3^-\) and eventually Cl\(^-\) in PM\(_{2.5}\) via neutralisation processes. The chemical composition data-base concerning PM\(_{2.5}\) (and PM\(_{10}\)), combined with predictions from ISORROPIA-II run in the forward mode, allow us to suggest that NH\(_3\) was most probably present in sufficiently high concentration to promote fine particle acidity neutralisation, during both the cold and the warm seasons. Although it is known that running ISORROPIA-II in the forward mode, with only aerosol concentrations as input may result in a bias in the predicted pH due to the repartitioning of ammonia in the model, this approach was the only one that allowed for a reasonable interpretation of the obtained results.

Over the warm seasons, ~ 35 % of the total analysed samples presented pH values in the very strong acidity fraction (0–3 pH units range), while over the cold seasons the fraction of samples in this pH range was ~ 43 %. However, while during the warm seasons ~ 24–25 % of the acidic samples were in the 1–2 pH range (reflecting mainly contributions from very strong inorganic acids), over the cold seasons a ~ 40 % fraction in the 1–3 pH range would reflect possible contributions from other acidic type species (i.e., organics). The observed changes in aerosols acidity could potentially impact the gas–particle partitioning of semi-volatile organic acids.
Acknowledgements

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Table 1: Basic statistics for the PM$_{10}$ and PM$_{2.5}$ fractions mass concentrations determined over the investigated period (n = 84 sampling events) in Iasi, north-eastern Romania.

<table>
<thead>
<tr>
<th>Statistical parameter</th>
<th>PM$_{10}$ (µg m$^{-3}$)</th>
<th>PM$_{2.5}$ (µg m$^{-3}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Working day</td>
<td>Weekend</td>
</tr>
<tr>
<td>Mean</td>
<td>19.25</td>
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<tr>
<td>Median</td>
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<tr>
<td>Stdev</td>
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</tr>
<tr>
<td>Min</td>
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<td>7.11</td>
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<tr>
<td>Max</td>
<td>42.65</td>
<td>44.84</td>
</tr>
</tbody>
</table>
Table 2: Annual and/or seasonal arithmetic means of the PM_{10} and PM_{2.5} fraction mass concentrations in Iasi, north-eastern Romania, and other various European sites (mean ± stdev).

<table>
<thead>
<tr>
<th>Site</th>
<th>Category</th>
<th>Sampling aagl (m)</th>
<th>Sampling period</th>
<th>PM_{2.5} (µg m^{-3})</th>
<th>PM_{10} (µg m^{-3})</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iasi (Romania) urban</td>
<td>35</td>
<td>2016</td>
<td></td>
<td>16.9±9.1</td>
<td>18.9±9.3</td>
<td>This work</td>
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<tr>
<td></td>
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<td>14.0±7.1 (warm)</td>
<td>16.8±8.3 (warm)</td>
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<td></td>
<td></td>
<td>21.3±13.0 (cold)</td>
<td>22.6±13.1 (cold)</td>
<td>Arsene et al., 2011a</td>
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<td></td>
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<td>15.6±8.7**</td>
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<td></td>
<td>9.1±5.5#</td>
<td>-</td>
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<tr>
<td>Iasi (Romania) urban</td>
<td>25</td>
<td>2007–2008</td>
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<td>10.5±11.2</td>
<td>38.3±25.4</td>
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<tr>
<td>Paris (France) urban</td>
<td>20</td>
<td>2009–2010</td>
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<td>14.8±9.6</td>
<td>-</td>
<td>Bressi et al., 2013</td>
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<td>Central western Europe (FR09)</td>
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<tr>
<td>Central Europe (DE44)</td>
<td>EMEP and 4 regional background sites</td>
<td>2012–2013</td>
<td></td>
<td></td>
<td></td>
<td>Alastuey et al., 2016b</td>
</tr>
<tr>
<td>Eastern Europe (SK06,HU02)</td>
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<tr>
<td>Eastern Europe (MD13)</td>
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<tr>
<td>South western Europe (ES22)</td>
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<tr>
<td>Central southern Europe (IT01)</td>
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<tr>
<td>South eastern Europe (GR02)</td>
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<tr>
<td>Thessaloniki (Greece) urban</td>
<td>7</td>
<td>2011–2012</td>
<td></td>
<td>37.7±15.7</td>
<td>-</td>
<td>Tolis et al., 2015</td>
</tr>
<tr>
<td>Thessaloniki (Greece) urban</td>
<td>3</td>
<td>2011–2012</td>
<td></td>
<td>21.5±8.3 (warm)</td>
<td>33.9±19.3 (cold)</td>
<td>Voutsa et al., 2014c</td>
</tr>
<tr>
<td>Finokalia (Greece) remote coastal</td>
<td>~ 5</td>
<td>2004–2006</td>
<td>18.2</td>
<td>30.8</td>
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<td>Gerasopoulos et al., 2007</td>
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<tr>
<td>Bologna (Italy) urban background</td>
<td>courtyard</td>
<td>2005–2006</td>
<td>31.6±21.0</td>
<td>44.5±24.2</td>
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<td>Tositti et al., 2014</td>
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<tr>
<td>Venice (Italy) semi-rural coastal</td>
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<td>2007–2008</td>
<td></td>
<td>22.5±12.9</td>
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<td>Masiol et al., 2012</td>
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<td>Prague (Czech) urban</td>
<td>12-25</td>
<td>2004–2005</td>
<td></td>
<td>33±13</td>
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<td>Schwarz et al., 2008</td>
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</table>

Note: *sampling altitude above ground level,**PM_{0.027-1.6} and *PM_{0.381-1.6} annual values given for discussion and comparison purposes with Arsene et al. (2011) study, *the total (coarse + fine) and the fine fractions reported (coarse fraction - particles of AED > 1.5 µm and fine fractions - particles of AED < 1.5 µm); **S–summer (8 June–12 July 2012), W–winter (11 January–8 February 2013); *averaged value of warm and cold seasons data (error propagation method for uncertainty estimation); SE12–Aspvreten (Sweden); IE321–Mace Head (Ireland); FR09–Revin (France); DE44–Melpitz (Germany), SK06–Starina (Slovak Republic), HU02–K-Puszta (Hungary), MD13–Leova II (Moldavian Republic), ES22–Montsec (Spain); IT01–Montelibretti (Italy), GR02–Finokalia (Greece).
Table 3: Monthly averages of meteorological variables, PM$_{10}$, PM$_{2.5}$ fractions and of water soluble ions mass concentrations (µg m$^{-3}$) in PM$_{2.5}$ aerosol particles from Iasi, north-eastern Romania. Data are presented as mean±stdev (median).

<table>
<thead>
<tr>
<th>Month</th>
<th>January</th>
<th>February</th>
<th>March</th>
<th>April</th>
<th>May</th>
<th>June</th>
<th>July</th>
<th>August</th>
<th>September</th>
<th>October</th>
<th>November</th>
<th>December</th>
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</thead>
<tbody>
<tr>
<td>WS (m/s)</td>
<td>6.41±4.89</td>
<td>7.34±4.79</td>
<td>6.49±4.79</td>
<td>6.41±5.05</td>
<td>5.45±4.44</td>
<td>6.07±4.36</td>
<td>6.28±5.00</td>
<td>5.37±4.72</td>
<td>4.67±7.33</td>
<td>7.17±4.57</td>
<td>5.72±9.42</td>
<td>3.77±3.76</td>
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<td>AT (°C)</td>
<td>-1.06±6.47</td>
<td>6.86±3.98</td>
<td>6.50±3.49</td>
<td>13.65±4.15</td>
<td>15.65±3.25</td>
<td>21.05±4.75</td>
<td>23.27±3.05</td>
<td>21.86±1.12</td>
<td>18.20±5.57</td>
<td>10.03±5.11</td>
<td>7.41±5.86</td>
<td>2.77±2.41</td>
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<td>RH (%)</td>
<td>71.27±15.86</td>
<td>68.20±16.97</td>
<td>46.30±27.50</td>
<td>44.43±13.19</td>
<td>50.32±15.88</td>
<td>46.63±16.69</td>
<td>32.95±7.58</td>
<td>37.12±13.14</td>
<td>31.61±14.10</td>
<td>54.30±10.88</td>
<td>69.92±22.10</td>
<td>81.85±13.22</td>
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<tr>
<td>n*</td>
<td>5</td>
<td>5</td>
<td>8</td>
<td>8</td>
<td>9</td>
<td>5</td>
<td>9</td>
<td>7</td>
<td>7</td>
<td>6</td>
<td>5</td>
<td>5</td>
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<tr>
<td>PM$_{2.5}$</td>
<td>23.39±11.65</td>
<td>21.30±9.73</td>
<td>16.10±5.31</td>
<td>15.29±8.34</td>
<td>8.98±3.78</td>
<td>11.45±3.61</td>
<td>16.15±11.12</td>
<td>14.00±4.27</td>
<td>17.36±5.60</td>
<td>12.71±4.81</td>
<td>16.93±13.09</td>
<td>30.94±9.51</td>
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<tr>
<td>PM$_{10}$</td>
<td>24.25±11.99</td>
<td>22.11±8.50</td>
<td>17.25±5.28</td>
<td>18.22±11.08</td>
<td>12.11±4.43</td>
<td>14.33±3.98</td>
<td>19.05±11.57</td>
<td>16.59±5.52</td>
<td>17.96±6.87</td>
<td>15.13±5.81</td>
<td>18.13±13.58</td>
<td>32.08±9.44</td>
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<tr>
<td>Cl$^-$</td>
<td>(0.34)</td>
<td>(0.37)</td>
<td>(0.10)</td>
<td>(0.15)</td>
<td>(0.13)</td>
<td>(0.14)</td>
<td>(0.18)</td>
<td>(0.31)</td>
<td>(0.18)</td>
<td>(0.32)</td>
<td>(0.19)</td>
<td>(0.64)</td>
</tr>
<tr>
<td>NO$_3^-$</td>
<td>3.54±1.93</td>
<td>3.21±1.36</td>
<td>2.42±1.09</td>
<td>1.36±0.86</td>
<td>0.47±0.28</td>
<td>0.31±0.17</td>
<td>0.41±0.17</td>
<td>0.69±0.41</td>
<td>1.25±0.82</td>
<td>1.88±1.27</td>
<td>3.62±1.10</td>
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<tr>
<td>SO$_4^{2-}$</td>
<td>2.76±1.66</td>
<td>2.58±0.91</td>
<td>2.03±0.71</td>
<td>1.96±0.75</td>
<td>1.70±0.81</td>
<td>2.39±1.31</td>
<td>2.04±0.98</td>
<td>2.22±0.76</td>
<td>2.15±0.84</td>
<td>1.96±1.00</td>
<td>1.75±0.27</td>
<td>2.16±0.69</td>
</tr>
<tr>
<td>CH$_3$COO$^-$</td>
<td>(0.45)</td>
<td>(0.63)</td>
<td>(0.29)</td>
<td>(0.82)</td>
<td>(0.49)</td>
<td>(0.57)</td>
<td>(0.48)</td>
<td>(0.57)</td>
<td>(0.60)</td>
<td>(0.89)</td>
<td>(0.43)</td>
<td>(0.54)</td>
</tr>
<tr>
<td>HCO$_3^-$</td>
<td>0.07±0.05</td>
<td>0.06±0.03</td>
<td>0.09±0.09</td>
<td>0.15±0.20</td>
<td>0.04±0.02</td>
<td>0.05±0.03</td>
<td>0.12±0.11</td>
<td>0.08±0.02</td>
<td>0.09±0.02</td>
<td>0.04±0.06</td>
<td>0.03±0.02</td>
<td>0.07±0.02</td>
</tr>
<tr>
<td>CO$_2$</td>
<td>0.08±0.06</td>
<td>0.08±0.04</td>
<td>0.08±0.04</td>
<td>0.08±0.05</td>
<td>0.06±0.04</td>
<td>0.09±0.06</td>
<td>0.12±0.09</td>
<td>0.14±0.05</td>
<td>0.10±0.08</td>
<td>0.06±0.04</td>
<td>0.02±0.02</td>
<td>0.10±0.04</td>
</tr>
<tr>
<td>HCO$_3^-$</td>
<td>0.39±0.12</td>
<td>0.48±0.24</td>
<td>0.39±0.20</td>
<td>0.65±0.67</td>
<td>0.32±0.17</td>
<td>0.64±0.56</td>
<td>0.51±0.24</td>
<td>0.70±0.26</td>
<td>0.80±0.28</td>
<td>0.46±0.33</td>
<td>0.59±0.12</td>
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</tr>
<tr>
<td>Na$^+$</td>
<td>0.14±0.05</td>
<td>0.17±0.08</td>
<td>0.12±0.08</td>
<td>0.41±0.31</td>
<td>0.19±0.14</td>
<td>0.13±0.05</td>
<td>0.13±0.08</td>
<td>0.18±0.09</td>
<td>0.14±0.09</td>
<td>0.25±0.16</td>
<td>0.12±0.15</td>
<td>0.17±0.09</td>
</tr>
<tr>
<td>NH$_4$</td>
<td>2.07±1.08</td>
<td>1.94±0.70</td>
<td>1.48±0.58</td>
<td>0.93±0.25</td>
<td>0.80±0.29</td>
<td>0.90±0.36</td>
<td>0.94±0.40</td>
<td>0.84±0.16</td>
<td>0.97±0.23</td>
<td>1.10±0.48</td>
<td>1.17±0.57</td>
<td>2.09±0.49</td>
</tr>
<tr>
<td>K$^+$</td>
<td>0.54±0.25</td>
<td>0.48±0.20</td>
<td>0.27±0.12</td>
<td>0.30±0.13</td>
<td>0.26±0.16</td>
<td>0.28±0.16</td>
<td>0.37±0.21</td>
<td>0.31±0.11</td>
<td>0.45±0.14</td>
<td>0.41±0.13</td>
<td>0.43±0.32</td>
<td>0.71±0.18</td>
</tr>
<tr>
<td>Mg$^{2+}$</td>
<td>0.03±0.01</td>
<td>0.03±0.02</td>
<td>0.04±0.02</td>
<td>0.01±0.01</td>
<td>0.00±0.01</td>
<td>0.02±0.01</td>
<td>0.00±0.01</td>
<td>0.05±0.04</td>
<td>0.02±0.01</td>
<td>0.03±0.01</td>
<td>0.01±0.01</td>
<td>0.03±0.01</td>
</tr>
<tr>
<td>Ca$^{2+}$</td>
<td>0.21±0.07</td>
<td>0.27±0.08</td>
<td>0.22±0.07</td>
<td>0.28±0.24</td>
<td>0.15±0.05</td>
<td>0.30±0.25</td>
<td>0.25±0.12</td>
<td>0.92±0.84</td>
<td>0.30±0.08</td>
<td>0.15±0.08</td>
<td>0.16±0.17</td>
<td>0.20±0.12</td>
</tr>
<tr>
<td>pH</td>
<td>4.20±2.32</td>
<td>4.17±2.53</td>
<td>3.94±2.81</td>
<td>3.88±2.97</td>
<td>3.95±3.10</td>
<td>4.14±2.92</td>
<td>4.58±2.87</td>
<td>5.06±2.73</td>
<td>4.24±3.19</td>
<td>4.25±2.75</td>
<td>3.86±2.55</td>
<td>4.12±2.16</td>
</tr>
</tbody>
</table>

Note: ‘n’ represent the number of aerosol sample events collected each month.
Table 4: Correlation matrix (Pearson coefficients) for major ionic species (Cl\(^-\), NO\(_3\)^-, SO\(_4\)^2-, CH\(_3\)COO\(^-\), HCOO\(^-\), C\(_2\)O\(_4\)^2-, HCO\(_3\)^-), Na\(^+\), NH\(_4\)\(^+\)(total), K\(^+\), Mg\(^{2+}\), Ca\(^{2+}\)) in fine aerosol particles from Iasi, north-eastern Romania, both for cold (a) and warm (b) seasons.

(a) PM\(_{2.5}\) (cold) Cl\(^-\) NO\(_3\)^- SO\(_4\)^2- CH\(_3\)COO\(^-\) HCOO\(^-\) C\(_2\)O\(_4\)^2- HCO\(_3\)^- Na\(^+\) NH\(_4\)\(^+\)(total) K\(^+\) Mg\(^{2+}\) Ca\(^{2+}\)

<table>
<thead>
<tr>
<th></th>
<th>Cl(^-)</th>
<th>NO(_3)^-</th>
<th>SO(_4)^2-</th>
<th>CH(_3)COO(^-)</th>
<th>HCOO(^-)</th>
<th>C(_2)O(_4)^2-</th>
<th>HCO(_3)^-</th>
<th>Na(^+)</th>
<th>NH(_4)(^+)(total)</th>
<th>K(^+)</th>
<th>Mg(^{2+})</th>
<th>Ca(^{2+})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cl(^-)</td>
<td>1.00</td>
<td>0.75</td>
<td>0.59</td>
<td>0.56</td>
<td>0.57</td>
<td>0.71</td>
<td>0.22</td>
<td>0.49</td>
<td>0.73</td>
<td>0.80</td>
<td>0.35</td>
<td>0.04</td>
</tr>
<tr>
<td>NO(_3)^-</td>
<td>1.00</td>
<td>0.91</td>
<td>0.37</td>
<td>0.74</td>
<td>0.87</td>
<td>0.13</td>
<td>0.04</td>
<td>0.98</td>
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<tr>
<td>SO(_4)^2-</td>
<td>1.00</td>
<td>0.43</td>
<td>0.72</td>
<td>0.87</td>
<td>0.05</td>
<td>0.02</td>
<td>0.96</td>
<td>0.72</td>
<td>0.07</td>
<td>0.11</td>
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<tr>
<td>CH(_3)COO(^-)</td>
<td>1.00</td>
<td>0.56</td>
<td>0.63</td>
<td>0.13</td>
<td>0.17</td>
<td>0.45</td>
<td>0.47</td>
<td>0.01</td>
<td>0.03</td>
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<tr>
<td>HCOO(^-)</td>
<td>1.00</td>
<td>0.85</td>
<td>0.04</td>
<td>0.01</td>
<td>0.76</td>
<td>0.65</td>
<td>0.12</td>
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<td>C(_2)O(_4)^2-</td>
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<td>0.89</td>
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<td>0.17</td>
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<td>HCO(_3)^-</td>
<td>1.00</td>
<td>0.56</td>
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<td>0.05</td>
<td>0.76</td>
<td>0.98</td>
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<tr>
<td>Na(^+)</td>
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<td>0.06</td>
<td>0.83</td>
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<tr>
<td>NH(_4)(^+)(total)</td>
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<td>0.08</td>
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<tr>
<td>K(^+)</td>
<td>1.00</td>
<td>0.08</td>
<td>0.09</td>
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<tr>
<td>Mg(^{2+})</td>
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</tr>
<tr>
<td>Ca(^{2+})</td>
<td>1.00</td>
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</tr>
</tbody>
</table>

(b) PM\(_{2.5}\) (warm) Cl\(^-\) NO\(_3\)^- SO\(_4\)^2- CH\(_3\)COO\(^-\) HCOO\(^-\) C\(_2\)O\(_4\)^2- HCO\(_3\)^- Na\(^+\) NH\(_4\)\(^+\)(total) K\(^+\) Mg\(^{2+}\) Ca\(^{2+}\)

<table>
<thead>
<tr>
<th></th>
<th>Cl(^-)</th>
<th>NO(_3)^-</th>
<th>SO(_4)^2-</th>
<th>CH(_3)COO(^-)</th>
<th>HCOO(^-)</th>
<th>C(_2)O(_4)^2-</th>
<th>HCO(_3)^-</th>
<th>Na(^+)</th>
<th>NH(_4)(^+)(total)</th>
<th>K(^+)</th>
<th>Mg(^{2+})</th>
<th>Ca(^{2+})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cl(^-)</td>
<td>1.00</td>
<td>0.57</td>
<td>0.10</td>
<td>0.81</td>
<td>0.37</td>
<td>0.08</td>
<td>0.81</td>
<td>0.87</td>
<td>0.02</td>
<td>0.76</td>
<td>0.83</td>
<td>0.63</td>
</tr>
<tr>
<td>NO(_3)^-</td>
<td>1.00</td>
<td>0.34</td>
<td>0.02</td>
<td>0.14</td>
<td>0.17</td>
<td>0.63</td>
<td>0.71</td>
<td>0.23</td>
<td>0.08</td>
<td>0.86</td>
<td>0.39</td>
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<tr>
<td>SO(_4)^2-</td>
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<td>0.01</td>
<td>0.66</td>
<td>0.76</td>
<td>0.17</td>
<td>0.05</td>
<td>0.97</td>
<td>0.43</td>
<td>0.03</td>
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<tr>
<td>CH(_3)COO(^-)</td>
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<td>0.11</td>
<td>0.30</td>
<td>0.07</td>
<td>0.11</td>
<td>0.55</td>
<td>0.05</td>
<td>0.08</td>
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<tr>
<td>HCOO(^-)</td>
<td>1.00</td>
<td>0.58</td>
<td>0.32</td>
<td>0.01</td>
<td>0.66</td>
<td>0.59</td>
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<td>0.01</td>
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<td>0.04</td>
<td>0.71</td>
<td>0.46</td>
<td>0.01</td>
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<td>0.45</td>
<td>0.06</td>
<td>0.13</td>
<td>0.12</td>
<td>0.99</td>
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<td>K(^+)</td>
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<td>Mg(^{2+})</td>
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<tr>
<td>Ca(^{2+})</td>
<td>1.00</td>
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FIGURE captions

FIGURE 1: Sectors contributions identified from classification of 2-day back trajectories of air masses ending at Iasi and representative backward trajectories of long and short range transport, Black Sea influence and African dust (shown trajectories correspond to sampling events).

FIGURE 2: Patterns of the monthly arithmetic mean concentrations and standard deviations in the PM10, PM2.5 and PM10/PM2.5 variables at Iasi, north-eastern Romania.

FIGURE 3: Size distribution histograms of aerosol particles mass concentration gravimetrically determined over both the cold and warm seasons.

FIGURE 4: Distribution of the aerosol pH predicted by ISORROPIA-II (forward mode) vs. the ion balance (a), sensitivity of aerosol pH predicted with the model to small changes in the input aerosol NH4+ concentration (b) and bar chart distribution in aerosol pH over the warm and cold season both for NH4+ derived from raw IC data (c) and NH4+ (total) (d).

FIGURE 5: Size distribution of averaged aerosol mass, NO3− SO42− and NH4+ concentrations over cold (a) and warm (b) seasons accompanied by the size distribution of ISORROPIA-II estimated pH and H+ mass concentration over both the cold (c) and the warm (d) seasons.

FIGURE 6: Seasonal variations for selected water-soluble ionic components in the PM2.5 fraction (a-h) and variation of the mixing layer depth at the investigated site (i). The inset distribution presented within NO3− seasonal variation reflects the contribution of the coarse fraction over the warm seasons. The horizontal black line represents the mean, the horizontal colored line – the median, the box – the 25–75% percentiles, the length of the whiskers plot – the 10 and 90% of observed concentrations, circles – outliers).

FIGURE 7: Regression analysis of the [NH4+] vs. (2×[SO42−]) (a, b), [NH4+] vs. ([NO3−] + 2×[SO42−]) (c,d) and [NH4+] vs. ([$\text{Cl}^-$] + [NO3−] + 2×[SO42−]) (e,f) dependences specific for PM2.5 particles.

FIGURE 8: The relative contributions, as monthly-based averages, of identified and quantified water soluble ions to total detected components in the 0.0276–0.0945 µm (a), 0.155–0.612 µm (b), 0.946–2.39 µm (c) and 3.99–9.94 µm (d) size range grouped fractions.

FIGURE 9: Size distributions of seasonal averaged mass concentrations for Cl−, NO3−, SO42−, NH4+ (a,b) and K+, Na+, Mg2+, Ca2+ (c,d) ions in atmospheric aerosols from Iasi, both during the cold and, respectively, the warm seasons.

FIGURE 10: Evidences of long range transport contributions from Saharan dust within the size distribution of particulate Na+, Ca2+, Mg2+, Cl− ions and aerosol mass (a) and of air masses buoyancy phenomena within the size distribution of particulate NH4+, NO3−, SO42−, Mg2+ ions and aerosol mass (b).
FIGURE 1

Long range transport

Black Sea influence

Short range transport

African dust
FIGURE 2

Period of the year

Jan Feb Mar Apr May Jun Jul Aug Sept Oct Nov Dec

µg m⁻³ (PM₁₀ and PM₂.₅ fractions)

PM₂.₅
PM₁₀
PM₂.₅/PM₁₀

PM₂.₅/PM₁₀

Period of the year
FIGURE 3

Dp (µm) 0.1 1.0 10.0
dM/dlogDp (µg m⁻³) 0 6 12 18 24 30 36
Cold seasons
Warm seasons

Dp (µm) 0.1 1.0 10.0
FIGURE 4

(a) pH (ISORROPIA-II - forward)

(b) pH total (ISORROPIA-II - forward)

\[ \text{pH}_{\text{total}} = 0.98 \times \text{pH}_{\text{raw}} + 0.15 \quad (r^2 = 0.99) \]

(c) pH interval

(d) pH interval

Relative contribution (%)
FIGURE 5

(a) Cold mass - gravimetrically determined

(b) Warm mass - gravimetrically determined

(c) Cold

(d) Warm

\( \frac{d M}{d \log D_p} + \sigma (\mu g m^{-3}) \)

\( d \text{NO}_3^-, \text{SO}_4^{2-}, \text{NH}_4^+ / d \log D_p + \sigma \)

\( d \text{H}^+ / d \log D_p + \sigma \)

\( \text{pH} \)

\( d \text{H}^+ \) from \( \text{NH}_4^+ \) (total)

\( \text{pH} \) from \( \text{NH}_4^+ \) (total)
FIGURE 6

(a) Cl (µg m⁻³) vs. Period of year
(b) NO₃ (µg m⁻³) vs. Period of year
(c) SO₄²⁻ (µg m⁻³) vs. Period of year
(d) C₂O₄²⁻ (µg m⁻³) vs. Period of year
(e) K⁺ (µg m⁻³) vs. Period of year
(f) NH₄⁺ total (µg m⁻³) vs. Period of year
(g) Na⁺ (µg m⁻³) vs. Period of year
(h) Ca²⁺ (µg m⁻³) vs. Period of year
(i) Mixing Layer Depth (m) vs. Period of year
FIGURE 7

2\[SO_4^{2-}\] (nmol m\(^{-3}\))

\[\text{[NH}_4^+\] = 1.76 \times (2\text{[SO}_4^{2-}\]) + 0.69 \quad (r^2 = 0.92)\]
\[\text{[NH}_4^+\] = 1.23 \times (2\text{[SO}_4^{2-}\]) + 0.76 \quad (r^2 = 0.94)\]

\[\text{[NH}_4^+\] = 1.02 \times (2\text{[SO}_4^{2-}\]) + 0.54 \quad (r^2 = 0.94)\]
\[\text{[NH}_4^+\] = 0.95 \times (2\text{[SO}_4^{2-}\]) + 0.49 \quad (r^2 = 0.96)\]

\[\text{[NH}_4^+\] = 0.99 \times (\text{[NO}_3^-\] + 2\text{[SO}_4^{2-}\]) + 0.11 \quad (r^2 = 0.98)\]
\[\text{[NH}_4^+\] = 0.95 \times (\text{[NO}_3^-\] + 2\text{[SO}_4^{2-}\]) + 0.14 \quad (r^2 = 0.96)\]

\[\text{[NH}_4^+\] = 0.68 \times (\text{[Cl}^-\] + \text{[NO}_3^-\] + 2\text{[SO}_4^{2-}\]) + 0.46 \quad (r^2 = 0.97)\]
\[\text{[NH}_4^+\] = 0.98 \times (\text{[Cl}^-\] + \text{[NO}_3^-\] + 2\text{[SO}_4^{2-}\]) + 0.06 \quad (r^2 = 0.95)\]
FIGURE 9

(a) Cold

(b) Warm

(c) Cold

(d) Warm

Dp (µm)

0.1 1 10

dCl-/dlogDp (µg m⁻³)

0.0

0.2

0.4

0.6

0.0

0.1

0.2

0.3

0.4

0.5

0.6

0.0

0.1

0.2

0.3

0.4

0.5

0.6

K+, Na+, Ca²⁺

Mg²⁺

0.0

0.1

0.2

0.3

0.4

0.5

0.6

0.0

0.1

0.2

0.3

0.4

0.5

0.6

K⁺

Na⁺

Mg²⁺

Ca²⁺
FIGURE 10

(a) Mass - gravimetrically determined

(b) April 2016

Gravimetrically determined

Mass - gravimetrically determined

NH\textsubscript{4}^+

Mg\textsuperscript{2+}

SO\textsubscript{4}^{2-}

NO\textsubscript{3}^{-}

Na\textsuperscript{+}

Ca\textsuperscript{2+}

Cl\textsuperscript{-}

mass - gravimetrically determined

Mass - gravimetrically determined

NH\textsubscript{4}^+

Mg\textsuperscript{2+}

SO\textsubscript{4}^{2-}

NO\textsubscript{3}^{-}

Na\textsuperscript{+}

Ca\textsuperscript{2+}

Cl\textsuperscript{-}

mass - gravimetrically determined

Mass - gravimetrically determined

NH\textsubscript{4}^+

Mg\textsuperscript{2+}

SO\textsubscript{4}^{2-}

NO\textsubscript{3}^{-}

Na\textsuperscript{+}

Ca\textsuperscript{2+}

Cl\textsuperscript{-}